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Fundamentals of  
SOIL SCIENCE



# Fundamentals of SOIL SCIENCE

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SECOND EDITION

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SECOND EDITION

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## PREFACE

*Fundamentals of Soil Science* was written primarily as a college textbook for use in the introductory or general course in soils. Also, it may be studied profitably by soil conservation technicians, agricultural agents, farmers, or anyone seeking a knowledge of the soil and of the principles underlying successful soil management.

The purpose of the book is fourfold: first, to give the reader the opportunity of becoming familiar with soils as natural units or entities and with their inherent characteristics; second, to develop in the student an understanding of the significance of fundamental soil properties; third, to set forth basic relationships between soils and plants; and, last, to give the reader an understanding of the principles involved in the practical use and conservation of the soil.

General principles of soil science are emphasized and explained in simple terms, technical discussions, particularly of debatable points, being omitted. Special attention is given to progress which has been made in fundamental principles since the publication of the first edition of this book. The material dealing with the chemical properties of soils has been expanded into a separate chapter. Likewise the subject matter concerned with soil material, formation, and classification has been altered and enlarged to include modern concepts and divided into three chapters in place of two. An entire chapter is devoted to the subject of soil conservation, and another to the soil resources of our country. Proven soil management practices are referred to, but the details of practices have been omitted to a large extent.

It has been necessary to go somewhat into the fields of chemistry, botany, geology, physics, microbiology, crop production, and agricultural engineering because soil science is closely related to them.

Chapter sequence in the book provides a logical treatment of the subject. The arrangement of the material in the chapters is designed to assist the student in an effective study of the subject matter presented. For example, the "objectives" call attention to the main subdivisions of the material under discussion, and the "questions" bring out the pertinent points in the discussion.

We take pleasure in expressing appreciation to Dr. C. H. Spurway, Professor J. O. Veatch, Dr. G. J. Bouyoucos, and Dr. A. H. Mick for

## Preface

assistance in the preparation of this book. In the revision of the text valuable help has been received from Professor W. T. McGeorge, University of Arizona, Dr. C. E. Kellogg, Bureau of Plant Industry, Soil and Agricultural Engineering, Dr. R. D. Hockensmith, Soil Conservation Service, Dr. H. A. Curtis, Tennessee Valley Authority, and Dr E. P. Whiteside, Dr. R. L. Cook, Dr. J. Q. Lynd, and other members of the Soil Science Department of Michigan State College.

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*East Lansing, Michigan*  
*July, 1951*

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# Materials for Soil Development

Soils are developed; they are not merely an accumulation of debris resulting from decay of rock and organic materials. Soil formation is, accordingly, a constructive as well as a destructive process. Destructive forces predominate in the breaking down chemically and physically of minerals and plant and animal structures, with the resultant partial loss of the more soluble and volatile products. Particles of the original materials in various stages of decomposition also remain, together with newly developed compounds. The constructive forces develop new chemical compounds, both mineral and organic, and provide a new distribution or association of the materials so that the resultant soil has a characteristic structural and textural as well as chemical composition which influences plant growth. In other words, a soil is an entity—an object in nature which has characteristics that distinguish it from all other objects in nature.

Soils are developed from two classes of materials: (1) partially decomposed plant substance (organic material); and (2) rocks and minerals (mineral material). To a limited extent soils develop in a mixture containing considerable quantities of both organic and mineral matter. In discussing the parent materials from which soils are developed it is helpful to consider the following objectives.

## Objectives

- A. Kinds and composition of mineral parent material.
- B. Characteristics of minerals from which soils are derived.
- C. Igneous rocks as parent material.
- D. Sediments as soil parent material.
- E. Nature and source of organic parent materials.

## KINDS AND COMPOSITION OF MINERAL PARENT MATERIAL

Before the forces of soil development can be operative, it is necessary that raw materials on which they can work be present. Let us

first study, therefore, parent materials from which soils are made. The mineral materials have been classified from a purely geological point of view, their mineralogical composition, texture, structure, and age being taken into account. Occasionally soil groups are incorrectly designated according to the geological classification of their parent material, and we carelessly speak of "granite soils" or "limestone soils." When we use these terms what we really mean is "soils developed from granite rocks" or "soils developed from limestone." In the consideration of mineral parent material, two questions suggest themselves.

### Questions

1. What is meant by "residual" and "transported" soils?
2. Which chemical elements make up a large part of the earth's surface?

**Residual and Transported Materials.** Soils have developed from the decay of hard rocks, consolidated sediments, or unconsolidated sediments exposed to soil-forming processes. Soils developed from hard rocks or consolidated sediments are often loosely termed residual soils. On the other hand, many soils have formed from stones, sand, silt, and clay which have been moved from their original sites and not consolidated after their deposition. Such soils are frequently called *transported soils*; more correctly speaking, they are soils that have developed in transported, unconsolidated material, because it is impossible for a soil to be naturally transported in its entirety. The general distribution of residual and transported soil materials in the United States is shown in Fig. 1.

**Chemical Composition of the Earth's Crust.** The solid crust of the earth (lithosphere) is made up largely of eight chemical elements. In fact, two elements, oxygen and silicon, compose some 75 per cent of it. On the other hand, many of the elements important in the growth of plants and animals occur in very small quantities. Needless to say, these elements and their compounds are not evenly distributed throughout the earth's surface. For example, in some places phosphorus compounds are so concentrated that they are mined, whereas in many other areas there is a deficiency of phosphorus for maximum plant growth. The composition of the lithosphere is shown in Fig. 3.

**Differences in Soil Parent Materials.** There were many variations in the original rocks from which soils were derived. Igneous, sedimentary, and metamorphic rocks all served as sources; and within these broad groups there are, of course, many differences in composition, structure, and ease of decomposition. A short discussion of igneous rocks and sedimentary materials will be given.

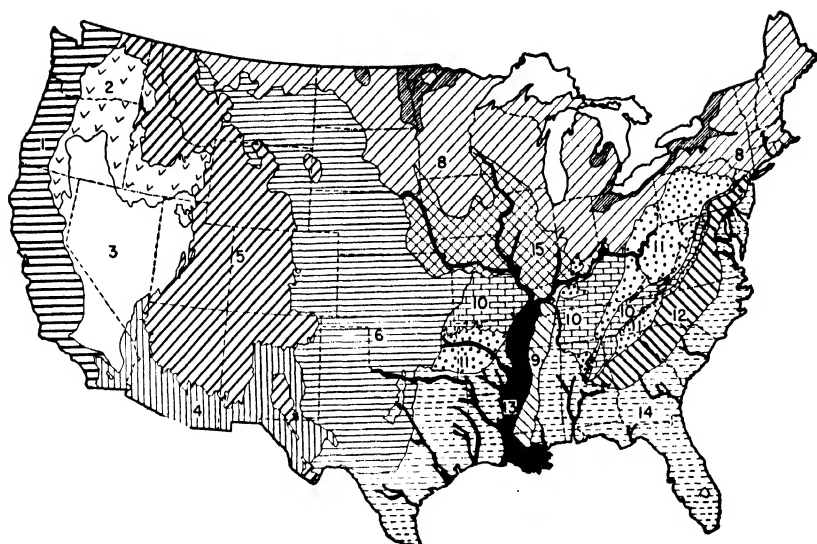


FIG. 1. A generalized physiographic map of the United States. [Drawn from a map prepared by the Division of Soil Survey and presented in *U.S.D.A. Bull. 96.*]

#### LEGEND OF AREAS

1. Pacific Coast region.
2. Northwest intermountain region.
3. Great Basin region.
4. Southwest arid region.
5. Rocky Mountain region.
6. Great Plains region.
7. Glacial lake and river terraces.
8. Glacial region.
9. Loessial deposits.
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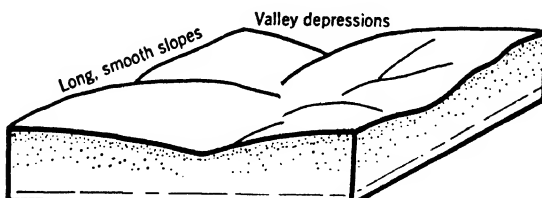


FIG. 2. A diagrammatic presentation of the mature topography of a residual soil area. The long smooth slopes and moderate valley depressions with well-developed drainages are characteristic. (Compare with Fig. 5.)

## Materials for Soil Development

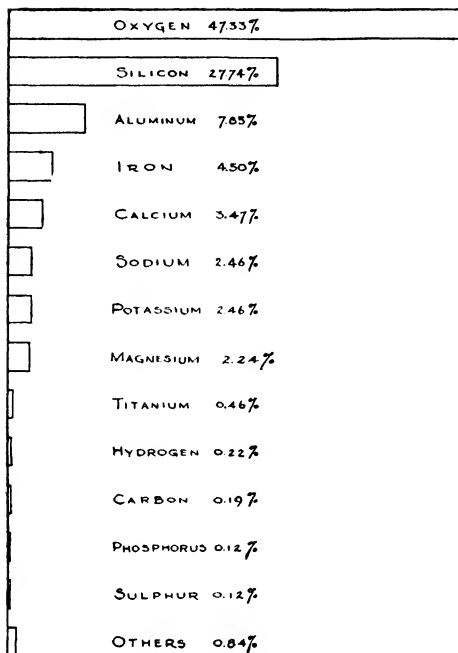


FIG. 3. Composition of the solid crust of the earth (lithosphere) to a depth of 10 miles. Note relatively small percentages of several elements essential for plant growth. [From F. W. Clarke, *U.S.G.S. Bull.* 695.]

### CHARACTERISTICS OF MINERALS FROM WHICH SOILS ARE DERIVED

Soils are composed very largely of fragments of rocks and minerals and of the products of their decomposition. A student should know at least some of the elementary facts concerning the composition and properties of minerals found most abundantly in soils. Three questions serve as guides in the study of soil minerals.

#### Questions

1. Minerals represent what kinds of chemical compounds?
2. Into what groups are soil minerals divided?
3. What are the chief characteristics of the mineral groups?

**Groups, Composition, and Characteristics of Minerals.** Combinations of oxygen with silicon and iron, and sometimes with water also, give rise to such compounds as  $\text{SiO}_2$  and  $\text{Fe}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$  and constitute one group of minerals called oxides. Basic compounds like  $\text{Ca}(\text{OH})_2$  and

## Groups, Composition, and Characteristics of Minerals 5

$\text{Mg}(\text{OH})_2$  combine with  $\text{CO}_2$  to form another mineral group, known as carbonates, which is very important in soil formation. By far the largest number of minerals belong to the silicate group, in which Ca, Mg, Na, K, Fe, and Al combine with the various silicic acids like orthosilicic ( $\text{H}_4\text{SiO}_4$ ), metasilicic ( $\text{H}_2\text{SiO}_3$ ), and polysilicic ( $\text{H}_4\text{Si}_3\text{O}_8$ ). At times the Al is part of the negative radical forming aluminosilicates. Many of these compounds and the products formed from their decomposition take on water; that is, they become hydrated.

The name, chemical composition, and some information about occurrence, properties, and decomposition products of some of the most important and widely distributed minerals and rocks are listed below.

### *Oxide Group*

|                             |  |   |
|-----------------------------|--|---|
| Quartz                      | $\text{SiO}_2$   | Very resistant to change; highly insoluble; very hard.  |
| Hematite                    | $\text{Fe}_2\text{O}_3$  | Red; hydrates with increase in volume to form $2\text{Fe}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ .  |
| Magnetite                   | $\text{Fe}_3\text{O}_4$  | Dark brown; magnetic, resistant to change; not of wide occurrence.  |
| Limonite                    | $2\text{Fe}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$   | Yellow; soft; resistant to change except in degree of hydration; widely distributed.  |
| Other hydrated oxides of Fe | $2\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$<br>$\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$<br>$\text{Fe}_2\text{O}_3 \cdot 2\text{H}_2\text{O}$ | Color varies from yellowish red to yellow with increase in hydration. Are supposed to account for red and yellow in soils. Are resistant to change other than in degree of hydration. Are of wide occurrence. |

### *Carbonate Group*

|           |   |  |
|-----------|---|--|
| Calcite   | $\text{CaCO}_3$ , quite pure.                     | Well crystallized. Goes into solution readily as $\text{Ca}(\text{HCO}_3)_2$ .   |
| Dolomite  | $\text{CaCO}_3 \cdot \text{MgCO}_3$               | Important constituent of dolomitic limestone.  |
| Magnesite | $\text{MgCO}_3$ , purity varies.                  | Not of wide occurrence.  |
| Siderite  | $\text{FeCO}_3$ , sometimes mined as an iron ore. | Not an important soil mineral. Found under poorly drained conditions. Combines with $\text{CO}_2$ and $\text{H}_2\text{O}$ to form $\text{Fe}(\text{HCO}_3)_2$ , which is soluble. |

*Feldspar Group*

Microcline and orthoclase



Orthoclase occurs in large quantities as a constituent of granite and gneiss.

Albite



Labradorite



Anorthite



The sodium and calcium feldspars form a part of many crystalline rocks like basalt, diabase, gabbro, diorite, and many lavas. The feldspars decompose fairly readily in the presence of  $\text{H}_2\text{O}$  and  $\text{CO}_2$ . The Ca goes to the carbonate or more soluble bicarbonate and the Na and K to carbonates. Some  $\text{SiO}_2$  is split off and may remain as insoluble  $\text{SiO}_2$  or form silicic acids and soluble or colloidal hydrated silicates. Most of the Al and much Si go into the formation of hydrated aluminosilicates, of which kaolinite ( $\text{H}_2\text{Al}_2\text{Si}_2\text{O}_8 \cdot \text{H}_2\text{O}$ ) is an example. The exact hydrated aluminosilicate formed depends on the conditions prevailing during decomposition. Compounds of this nature are important constituents of clay and play prominent roles in determining the properties of soils and the reactions that take place in them. Calcium feldspars decompose much more easily than do sodium and potassium feldspars.

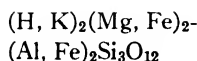
*Micas*

Muscovite  
(white)



Break up into fine glittering scales which are sometimes mistaken for gold when yellow. When much iron is present in biotite, it decomposes readily; otherwise the micas break up physically quite easily but de-

Biotite (dark)



compose chemically very slowly. They are abundant ingredients of many rocks.

### *Other Silicates*

|                                  |  |   |
|----------------------------------|--|---|
| Amphiboles<br>(hornblende)       | $\text{Ca}(\text{Mg}, \text{Fe})_2\text{Si}_4\text{O}_{12}$<br>with $\text{Na}_2\text{Al}_2\text{Si}_4\text{O}_{12}$ and<br>$(\text{Mg}, \text{Fe})_2(\text{Al}, \text{Fe})_2\text{Si}_2\text{O}_{12}$                   | The minerals are dark green in thin sections but appear black as they occur. Hornblende decomposes fairly readily because of its easy cleavage and high content of ferrous iron. On decomposition it produces a rust-colored clay which is a constituent of many red soils. Augite does not cleave so easily and often occurs as "black gravel" in soils. |
| Pyroxenes<br>(augite)            | $\text{CaMgSi}_2\text{O}_6$ with<br>$(\text{Mg}, \text{Fe})(\text{Al}, \text{Fe})_2\text{Si}_2\text{O}_6$<br>There is considerable variation in the proportions of the different elements, especially of Ca, Mg, Al, Fe. |   |
| Olivine                          | $\text{MgFeSiO}_4$   | Decomposes in presence of moisture to form serpentine ( $\text{H}_4\text{Mg}_3\text{Si}_2\text{O}_9$ ), liberating $\text{SiO}_2$ and $\text{FeO}$ .  |
| Leucite                          | $\text{KAlSi}_3\text{O}_{12}$  | A constituent of many volcanic rocks but otherwise not an abundant mineral. It is of interest because of its high K content.  |
| Talc (soap-stone) and Serpentine | $3\text{MgO} \cdot 4\text{SiO}_2 \cdot \text{H}_2\text{O}$<br>$3\text{MgO} \cdot 2\text{SiO}_2 \cdot 2\text{H}_2\text{O}$  | These two hydrated silicates of magnesium are found extensively in some regions. Both are soft, but serpentine in the absence of definite structure disintegrates slowly unless considerable ferrous iron is present. Talc cleaves readily and is very soft; so disintegration is accelerated. Both give rise to rather infertile soils.                  |

### *Sulfides*

|                            |                |   |
|----------------------------|----------------|---|
| Pyrite<br>and<br>Marcasite | $\text{FeS}_2$ | These minerals are yellow, often mistaken for gold when small fragments are mixed with sand or gravel. They oxidize under suitable conditions, forming $\text{H}_2\text{SO}_4$ . They are common constituents of rocks in some areas. |
|----------------------------|----------------|---|

*Phosphates*

|         |                                      |   |
|---------|--------------------------------------|---|
| Apatite | $[Ca_3(PO_4)_2]_3 \cdot Ca(F, Cl)_2$ | A dark brown mineral widely distributed but in small quantities. It is the original source of the phosphorus found in other combinations in the soil. |
|---------|--------------------------------------|---|

## IGNEOUS ROCKS AS PARENT MATERIAL

Igneous rocks were formed from a molten mass which was submitted to more or less rapid cooling conditions. With slow cooling, deep in the earth, large crystals of the mineral components formed, giving rise to the intrusive or plutonic rock types. Rapid cooling near or on the earth's surface resulted in small crystals or else in a glassy mass. These rocks constitute the extrusive or eruptive types. The mineral compounds in igneous rocks had no opportunity to be acted upon by weathering agencies before they were exposed at the earth's surface. As a result, many of the chemical combinations are very unstable and are easily broken down into more change-resistant compounds. The decay of igneous rocks may produce residues either high in calcium, magnesium, and other basic elements or low in such constituents and high in acidic, siliceous compounds. Some igneous rocks undergo rapid physical disintegration to produce sandy or gravelly residues, whereas other kinds decay into silt and clay, thus giving rise to fine-textured soils.

There are, then, many wide differences in soil parent materials of the igneous group. These differences influence to a greater or lesser extent the characteristics of the soils that develop in them. Great differences and extreme variations are, therefore, common in soils developed from these materials. A brief discussion of igneous rocks may be helpful.

*Igneous Rocks*

|                          |  |  |
|--------------------------|--|--|
| Granites<br>(intrusive)  | Chief mineral constituents are quartz, orthoclase, and occasionally plagioclase.   | Members of this group are the most acid; that is, they contain the highest percentages of silica of all igneous rocks. The mineral crystals may be very small or several inches in length in different kinds of granites. In the rhyolites they are small. Color varies from gray to shades of green, yellow, pink, and red. These are among the most common of igneous rocks. |
| Rhyolites<br>(extrusive) | One or more minerals of the mica, hornblende, or pyroxene group are usually present with minute particles of several other minerals. |  |

|                          |  |   |
|--------------------------|--|---|
| Syenites<br>(intrusive)  | Differ from granites essentially in the absence of quartz.   | Syenites and trachytes are intermediate in acidity between the granites and basalts.  |
| Trachytes<br>(extrusive) |  |   |
| Diorites<br>(intrusive)  | Differ from previous groups in the absence of potash minerals like orthoclase.                                 | Diorites and andesites are of wide distribution. Color varies from green to almost black.   |
| Andesites<br>(extrusive) | Principal constituents are plagioclase and hornblende.   |   |
| Gabbros<br>(intrusive)   | Gabbros and basalts contain primarily an Na·Ca feldspar, one or more pyroxenes, and sometimes olivine.         | These rocks contain less SiO <sub>2</sub> and more Na, Ca, and Mg than members of preceding groups. They are known as basic rocks. Because of their low silica content and high content of basic elements, the products of decomposition are basic. They are dark, ranging from gray or greenish to black, with browns and reds. Basalts are widely distributed among the younger volcanic rocks. In the United States they are found mainly west of the Mississippi. |
| Basalts<br>(extrusive)   | Apatite, mica, and hornblende are common constituents.   |   |
| Glasses                  | These rocks cooled so rapidly that there was no opportunity for crystallization, and hence they are amorphous. | Similar to granite in chemical composition.   |

### SEDIMENTS AS SOIL PARENT MATERIAL

As the name implies, sediments are materials which were deposited after transportation. They may be divided into two general groups on the basis of solidification after deposition, namely, (1) sediments which have been hardened into rock-like formations such as limestone, sandstone, and shale, and (2) relatively unconsolidated materials such as stones, sand, silt and clay and mixtures of them. It is customary to separate unconsolidated sediments into three groups on the basis of the agency which transported them, as wind, water, or ice. The materials are designated respectively as aeolian, alluvial, and glacial, and they will be discussed under these headings.

**Questions**

1. Where are glacial deposits found in the United States?
2. What land forms were deposited by glaciers?
3. What effect did glaciation have on topography?
4. Were there differences in the composition of glacial till?
5. Where are alluvial materials deposited?
6. What physical forms and characteristics do alluvial deposits exhibit?
7. What are the origin and characteristics of the coastal plains?
8. The hardened or rock-like sediments may be divided into what groups?
9. Where are the main loessial deposits in the United States?
10. What are some of the characteristics of loess?

**Glacial Materials.** Ice was the transporting agency for most of the surface mantle in northeastern United States and in the upper Mississippi Valley. The Ohio and Missouri rivers form a general southern

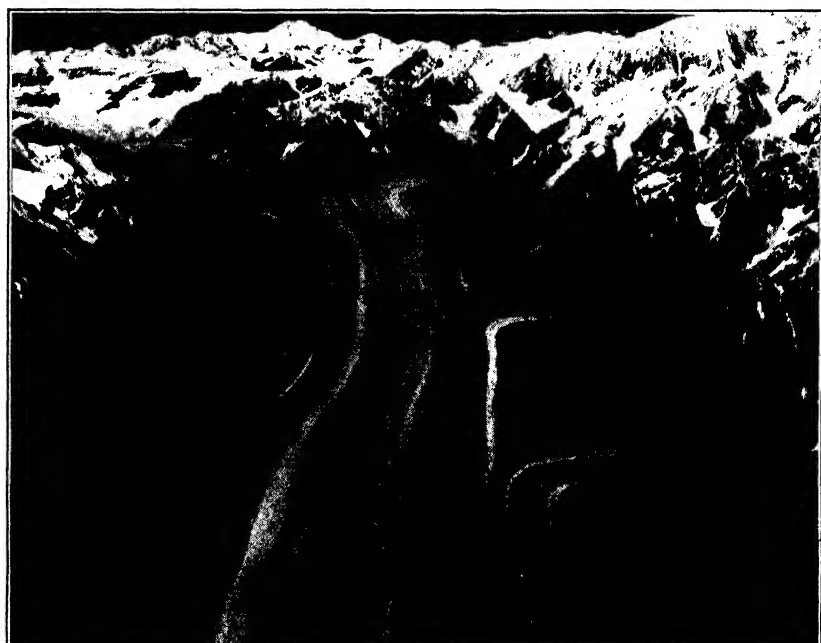


FIG. 4. The origin of a glacier. Notice the parent snow fields in the background and the small tributary glaciers. [Photograph by Bradford Washburn.]

boundary for ice-carried material. As the great continental ice-sheets moved southward from their accumulation centers in the Canadian

plateaus, they first followed and filled the great drainage valleys and then gradually spread out over the intervening upland and divides. As the ponderous ice-mass moved forward, it pushed before it and gathered within itself a large part of the unconsolidated surface layer. It also scooped up great rock fragments which scraped at the rock floor over which they passed. Sharp corners and edges of even the hardest rocks were ground smooth by this abrasive action to form the rounded rocks and boulders that are characteristics of glaciated landscapes. Large quantities of weathered and unweathered rocks, varying in size from

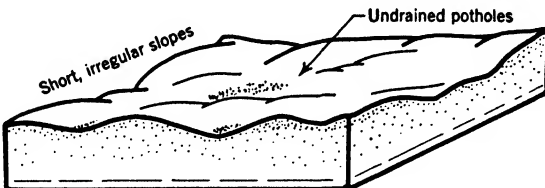


FIG. 5. Recently glaciated areas are geologically young and present a landscape with short, irregular slopes and small, poorly drained depressions known as "potholes." (Compare with Fig. 2.)

fine rock powder to massive boulders, were thus incorporated into the ice and carried along in the glacier.

**Land Forms Produced by Glaciers.** The movements of this continental ice-sheet depended on the changes in climatic conditions which took place during the glacial age. During mild periods the ice melted rapidly. In cold seasons melting ceased and the ice front would creep southward. Sometimes during extremely mild periods the ice would melt faster than it was pushed forward. This would lead to a rapid recession of the ice front, and all debris carried in the ice was, of course, dropped. Generally, after this type of recession, the land surface appeared as a rolling plain, called a *till plain* or *ground moraine*. At certain times climatic conditions allowed the glacier to melt back just as fast as its rate of advance, and this process resulted in the front of the ice remaining at one place for some time. All debris carried by the ice was brought to the line of the stationary ice-front and there dumped as melting proceeded. This process resulted in the formation of ridges or a series of hills, called *terminal* and *recreational moraines*. Moraines are usually composed of an unassorted, heterogeneous mass of boulders, rocks, sand, silt, and clay, briefly called *till*, but in places a water-sorting also occurred. As would be expected, the proportions of these materials vary greatly, and hence some moraines are

relatively high in sand content whereas others contain a large proportion of fine particles. Not only does the material vary in texture, but the shape of the surface is also variable.

As the ice melted, giving rise to moraines, great volumes of water rushed away. These waters carried quantities of sediment, the coarser of which was deposited as the current diminished. These coarse-textured, comparatively level deposits are known as *outwash* plains. They

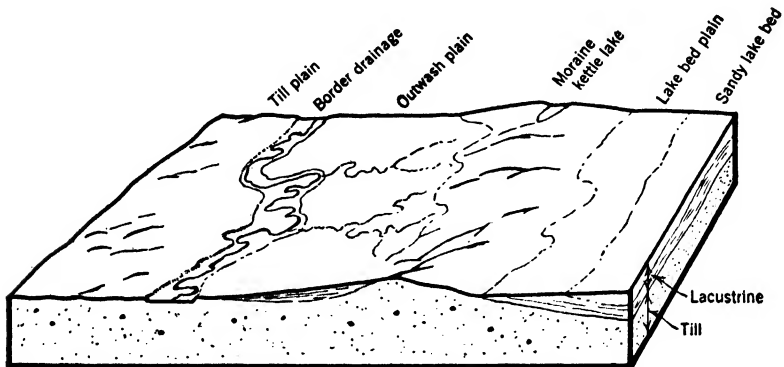


FIG. 6. A generalized view of physiographic features in a glaciated area. On the left is a till plain separated from the moraine by an outwash plain and a border-drainage way developed while the ice-front was at the moraine line and melting as rapidly as it advanced to deposit the morainic material. To the right a glacial lake has receded to leave a plain of heavy sediment partially covered with lacustrine sands (deposit nearer the water's edge).

will be discussed more fully in the sections dealing with alluvial materials. Most of the finer silt and clay were carried into slowly moving water or lake basins, where they settled out to form lake bed or *lacustrine* plains. In Michigan there is evidence of as many lakes which have become extinct as there are lakes in existence, whereas along the Great Lakes extensive areas have been exposed through the disappearance of ice barriers, lowering of the outlet at Niagara Falls, and tilting of the land surface. The disappearance of Lake Agassiz has laid bare a great land surface in Minnesota and the Dakotas, and the basin of old Lake Bonneville occupies an immense area in Utah.

The glaciers also produced *kames* and *eskers*, but these formations are only of local significance. Minor readvances of the ice-sheet resulted in the modification of these glacial formations in many places. Sometimes till was relaid over a lake plain, or sandy and gravelly outwash was pushed up to form gravelly moraines. Major readvances

"streamlined" glacial material in some parts of the country to form *drumlins*. It is customary to designate all the material deposited by glaciers and their melt waters as *glacial drift*. In general, the effect of glaciation was to scrape off and smooth the tops of high land forms while filling valleys and depressions. Thus continental glaciation decreased the local relief intervals and flattened topography.

**Differences in the Composition of Glacial Till.** The *chemical composition* of the debris carried by ice-sheets depends on the nature of the rocks over which they passed. Glaciers that originated near Hudson Bay gathered grayish till as they moved southward. Glaciers that came from north and west of Lake Superior gathered boulder clay that was high in iron-bearing minerals and red in color. Glacial formations, such as moraines, till plains, and lake deposits, produced by these two different glaciers therefore show many differences in chemical composition. In some places these two kinds of till, even though from distant sources, are closely associated; red materials, for example, may be directly on top of the gray drift of an earlier ice invasion.

The chemical composition of glacial drift often was modified by being mixed with an earlier, more weathered material. Although some portions were covered by only one ice-sheet, most of the glaciated area was invaded at least twice and in places as many as five times by major continental glaciers. Between invasions there elapsed long periods of mild weather during which soils developed in the glacial till and vegetation flourished. The next advance of the glacier scraped off these soils and redeposited them with other fresh debris. The old material, crushed and ground anew and mixed with rock powder of more recent origin, supplied a large amount of fresh, undecayed mineral surface, potentially rich in plant nutrients. The resulting till, however, differed from the old drift as well as from the new unmixed boulder clay and thus introduced another variation in the chemical differences of glacial materials in general.

Within the area covered by till from a single source there is, moreover, considerable variation in *physical* as well as chemical properties. Indeed, chemical and physical differences are closely associated. The chemical differences may originate in the mechanical separation by sedimentation of the various mineral components of boulder clay because of their different physical characteristics. For example, rocks containing a high proportion of quartz are hard and do not break down so readily as rocks containing more calcium, magnesium, or iron. It follows that sands and gravels, being the coarse components of drift, are

frequently the siliceous components. Clays, on the other hand, are the least siliceous part of boulder clay. When glacial streams, then, segregate the sand and gravel in outwash plains and the clays in lacustrine plains, they are at the same time separating the siliceous minerals from the basic minerals. However, even within each size group of materials there is variation in the content of limestone and soft minerals.

Glacial materials vary, then, not only in texture but also in chemical and mineralogical composition. This variation is due in part to the nature of the sources of the material. Other factors furthering this difference are the method of transportation, the many readvances of the continental glaciers, and finally the reworking and sorting by glacial streams. A common characteristic of all glacial parent materials is the extreme variability which naturally leads to local differences in the soils which develop from them.

**Alluvial Materials.** Alluvial parent materials are scattered in narrow, irregular strips bordering streams and rivers throughout glacial as well as residual areas. The largest uniform body of alluvium is located in the southern Mississippi Valley. Alluvial material ranges in texture from fine clay to large rocks and even to boulders. The fine sediments are washed into the drainage valley from the watershed surface. When the current is swift, the carrying capacity of the stream is high, but as the rate of flow decreases first the coarse and then the fine materials settle out. A common characteristic of all alluvial material is stratification, layers of different-sized particles overlying each other.

Most alluvium is carried and deposited during floods because it is at this period that erosion is most active and the carrying capacity of streams is at a maximum. When a flooding stream overflows its banks, its carrying power is suddenly reduced as the flow area increases and velocity decreases. This causes the coarse sands and gravels to settle along the bank where they sometimes form conspicuous ridges called *natural levees*. As the water reaches the *flood plains* of the valley, the rate of flow is slow enough to permit the silt to settle. Finally the water is left in quiet pools, from which it seeps away or evaporates leaving the fine clay. Levees are characterized by good internal drainage during periods of low water, whereas flood plains exhibit poor internal drainage.

Large mature valleys like the Mississippi are characterized by rivers with many meanders which develop in a well-recognized cycle. As a stream swings back and forth across its flood plain, cutting new channels in flood periods, sections of the old bed, known as *ox-bows*, are left as

lakes and swamps which fill with sediments during subsequent overflows.

*Terraces* are developed from flood plains as streams cut deeper channels because of lowered outlets. Several terraces may be found along a stream or a lake which has undergone repeated changes in level. Extensive terraces were formed as the glaciers receded and their outwash plains were no longer covered with water. Terraces usually are quite well drained and may be droughty. They exhibit stratification.

Streams flowing from hills or mountains into dry valleys or basins drop their sediments in a fan-like deposit as the water spreads out. These *alluvial fans* are usually coarse-textured, being composed of sands and gravels, and are well or excessively drained.

Sediments not deposited as flood plains are carried to the lake, gulf, or other body of water into which a stream empties. The decrease in velocity at the stream's mouth together with the coagulating effect of the salt content of the receiving water body results in the deposition of much of the suspended material, thus producing a delta. These deposits are poorly drained but, where drainage is provided, they constitute important crop-producing areas, as is evidenced by the deltas of the Nile, Po, Tigris, Euphrates, and the Mississippi.

Flood plains as well as deltas are in general rich in plant nutrients and comparatively high in organic matter content. Terraces and alluvial fans, on the other hand, are more likely to be less fertile. Special crops such as vegetables and fruits frequently are grown on the latter formations because the soil warms up quickly and their good drainage and coarse texture permit free root development.

Along much of our Atlantic Coast and extending around the Gulf (see 14 in Fig. 1) is a large area of land known as the Coastal Plains. This material was derived from sediments carried by streams and deposited in the ocean and gulf through decreased current velocity and chemical coagulation. Much of the area is sandy but is interspersed with beds of silt and clay which were deposited in estuaries or other sheltered bodies of water or farther out in the ocean. When raised above water level these deposits were subjected to soil-forming processes.

The solvent power of ocean water, coupled with the ceaseless grinding by wave action, pulverized soft minerals and extracted soluble salts. As a result large areas are quite sandy, composed mainly of particles of hard minerals, and are low in plant-nutrient content. Nevertheless a large acreage is used for the production of vegetables, fruits, and other special crops through the use of liberal quantities of fertilizer. There are also

extensive areas of forest. Considerable acreages of loam, silt loam, and clay loam, especially in the coastal plains of Texas and Louisiana, are used for general farming and production of special crops.

**Sedimentary Rocks.** Large areas of sedimentary rocks are composed of materials deposited in relatively large bodies of water such as lakes and oceans. The sediments are derived from the disintegration and decomposition of rocks and minerals on the earth's surface. The material is transported by water in suspension and in solution. The coarser particles such as sand are deposited near the shore line, but the silt and clay are carried out farther and settle out more slowly. Some of the materials in solution, primarily the bicarbonates of calcium and magnesium, are transformed into the less soluble carbonates by chemical processes, by shell-forming animals, and by growth of certain plants in the water and so deposited. By chemical and geological processes some of these sediments are solidified into rock; some are cemented quite firmly, and others are weakly cemented or remain unsolidified. Ultimately, large areas of them are elevated by geological forces, become a part of the earth's surface, and serve as parent material for soil formation. Here is presented a brief grouping and description of sedimentary rocks which have been more or less solidified. This list does not, of course, contain all the groups of sedimentary rocks, as beds of clay, glacial till, wind deposits, etc., are so classed by geologists.

|                |  |  |
|----------------|--|--|
| Limestone      | $\text{CaCO}_3$ ; varies greatly in purity. Main impurities are $\text{Fe}_2\text{O}_3$ , $\text{SiO}_2$ , $\text{MgCO}_3$ , clay, and organic matter. | Dense and compact but relatively soft. Goes into solution readily as $\text{Ca}(\text{HCO}_3)_2$ , leaving impurities. Of wide occurrence and great importance in soil formation.                        |
| Dolomite       | $\text{CaMg}(\text{CO}_3)_2$ ; composition varies within limits. Impurities as in limestone.   | Harder than limestone, and decomposes with greater difficulty. Goes into solution slowly as $\text{Ca}(\text{HCO}_3)_2$ and $\text{Mg}(\text{HCO}_3)_2$ , leaving impurities. Rather widely distributed. |
| Marl and chalk | $\text{CaCO}_3$ ; impurities are clay, organic matter, sand, $\text{MgCO}_3$ . Purity varies widely.   | An unconsolidated to somewhat solidified form of limestone.  |
| Shales         | Composed mainly of feldspars, clays, minerals, and quartz.   | Represents clays which have been solidified to different degrees of hardness.  |

|           |   |   |
|-----------|---|---|
| Sandstone | High in content of quartz. Contains smaller quantities of other minerals. May be cemented with silica, $\text{CaCO}_3$ , and iron compounds. From none to few per cent of $\text{CaCO}_3$ . | Large areas. Important in soil formation. |
|-----------|---|---|

It should be remembered that sedimentary rocks are composed of the fragments of igneous rocks and their decomposition products, which were transported, deposited, and more or less solidified. They exhibit a great variation in composition. Sandstones usually are high in content of quartz and other minerals rich in silica. The cementing materials frequently are compounds of iron with a calcium carbonate content varying from zero to a high percentage. Clay or silt sediments have been converted into shale and slate, which may contain either no calcium carbonate or a considerable amount. Some limestones are mainly calcium carbonate in the crystalline form of calcite, whereas others may contain varying proportions of dolomite and are termed dolomitic limestones and dolomites. Some limestone also contains appreciable quantities of phosphorus compounds, as in the Bluegrass region of Kentucky, parts of Tennessee, and in Florida. The soils formed by these phosphatic limestones are rich in phosphorus although they may be acid through leaching away of the lime.

**Wind-Transported or Aeolian Soil Material.** There are three distinct classes of wind-moved soil material: (1) sand of variable fineness which may be collected into low swells or steep ridges, like the dunes occurring on the leeward side of large bodies of water and on sandy deserts; (2) volcanic ash, deposits of which are found in Kansas, Nebraska, and Montana; and (3) silt-like material, called *loess*, which occupies large areas in the United States, Europe, and Asia.

*Dune sand* is of little agricultural value, although crops are produced on it to a limited extent in humid regions. At times dunes are a hazard to agriculture, for in their movement they sometimes gradually cover good land.

Soils derived from *volcanic ash* are of little agricultural value in the United States. They are very porous and light in weight.

*Loess* was deposited in central United States after the recession of the ice-sheet. This material was derived in part from sediments deposited by huge rivers which were fed by the melting continental glaciers in a broad belt, even beyond the southern limits of glaciation. A period of aridity after the recession of the glaciers with strong westerly winds set the

stage for the transportation of this wind-blown material to its present resting place, as shown in Fig. 7. The great thickness of the deposits of loess on the east and northeast banks of the Mississippi and Missouri rivers is one of the facts which has led to this explanation of the accumulation of the material. Glaciers from the Rockies probably supplied the sediments making up the western part of the loessial deposits.

Extensive deposits of loess are also found along the Rhine and its



FIG. 7. Loess was being deposited when this picture was taken. During this Colorado dust storm total darkness lasted for half an hour. [Courtesy of Soil Conservation Service.]

tributaries and over a large part of the immense valley of the Hwang Ho. Other deposits occur in southern Russia, several of the Balkan countries, and northern France, Belgium, and Poland.

Loess is composed largely of silt and is commonly grayish yellow or buff. This wind-blown material has the property of standing in almost vertical walls so that gullies and streams cutting through it have very steep banks. Its content of mineral plant nutrients was originally high, as was the quantity of calcium compounds. Loess is one of the most uniform of soil parent materials, but even it varies considerably in texture and mineralogical composition. Soils developed from these deposits are frequently referred to as fertile; however, no single material always gives rise to productive soil as the parent material is only one of the factors involved in soil formation.



FIG. 8. Loessial material has the characteristic of standing in almost vertical cliffs (a roadside cut near Natchez, Mississippi).

#### NATURE AND SOURCE OF ORGANIC PARENT MATERIALS

In locations where considerable quantities of plant material grow and decay is limited because of much water or low temperatures, a large accumulation of partially decayed vegetable matter gradually develops. Such deposits are of wide occurrence and are not restricted to any given climatic zone. They are found in Europe, Asia, Africa, Canada, South America, and United States, and various other places including the tropics. It may be said, however, that accumulations of this nature are more common in northern latitudes and occupy a larger percentage of the land surface in Norway, Sweden, Ireland, Scotland, northern Germany and Russia, and Holland than in countries lying farther south. In the Tundra region organic deposits are of frequent and extensive occurrence. Because of variation in chemical and physical characteristics and in location, numerous terms have been used to designate these deposits. As a matter of convenience in this discussion they will be re-

ferred to as peat. Several questions will emphasize the principal characteristics and facts relative to formation of these accumulations.

### Questions

1. What is the extent and distribution of peat in the United States?
2. Under what conditions is peat accumulated in this country?
3. Is there considerable variation in the material making up a peat deposit?

**Area of Peat Not Accurately Determined.** No accurate estimates have been made of the acreage of peat in the United States. Figures given in various publications concerning this point vary greatly. One difficulty encountered in making a dependable estimate of peat acreage is the inability to distinguish between areas of swamp, mineral soil, and organic soil deposits without a rather careful survey. Perhaps it is safe to say that there are 20,000,000 to 30,000,000 acres of organic soil in continental United States, excluding Alaska. Although the area of peat is not great in comparison with the total land surface of our country, the land rates high in terms of value of crops produced per acre when it is properly developed and managed. It should be remembered also that only a small part of the peat land in this country is producing crops.

The distribution of peat lands presents an interesting pattern. Florida and Minnesota have the largest areas, with possibly 5 to 6 million acres each. Michigan comes next with 4 to 6 million acres. Wisconsin contains a large acreage, possibly 3 million, and areas of appreciable extent occur in Maine, New Jersey, New York, Ohio, Indiana, Virginia, and several other states.

**Slow Decay Essential for Peat Production.** Even when a rank growth of vegetation occurs each year and remains on the soil, peat does not accumulate unless decay processes are very slow. The most common factor which limits decay is an excess of water. Accordingly in moderate to warm climates peat accumulates in shallow lakes and swamps. The topography, resulting from glaciation, in Minnesota, Wisconsin, Michigan and to a less extent in Maine, New York, and New Jersey, has given rise to innumerable small lakes and swamps. Thus conditions have been suitable for peat accumulation. Along the Atlantic and Gulf Coast many swamps have developed because of the slight elevation above sea level. Peat has accumulated in many of these.

In cold climates with short summers but very long days, vegetation grows very rapidly. Also types of plants, such as mosses and lichens, which grow in relatively low temperatures are abundant. Because of the long winters in such areas, decay of vegetable matter is restricted and accumulations of organic soil material develop.

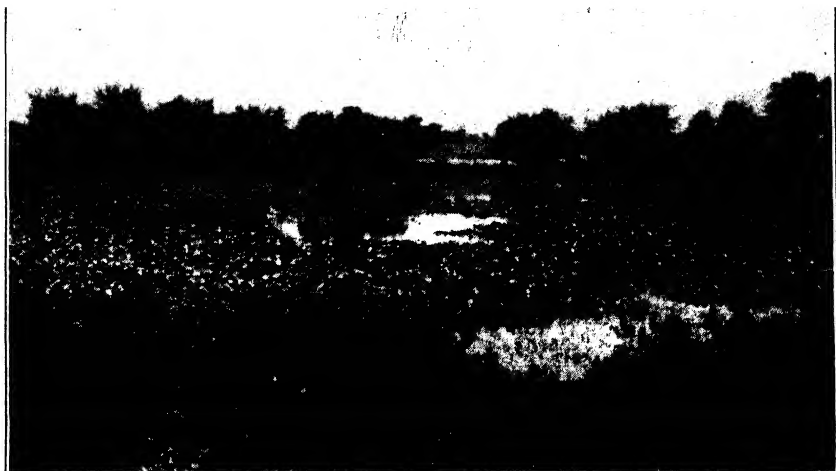


FIG. 9. A lake which is rapidly being filled with vegetation and will soon become a peat swamp.

**Great Variation in Characteristics of Organic Soil Material.** The characteristics of organic soil materials depend largely on the nature of the vegetation from which they developed. For example, in relatively deep water remains of algae and other aquatic plants give rise to highly colloidal material which shrinks greatly on drying. As the lake gradually fills, rushes, wild rice, water lilies, and similar plants flourish. The partially decayed remains of these plants are less slimy and colloidal. Gradually sedges, reeds, and eventually grasses are able to grow. Peat from such plants is much more fibrous than that produced from plants growing in deeper water. Shrubs and trees follow in time and produce a woody type of peat. Changes in water depth may cause a recurrence of deeper-water plants, and hence layers of more pulpy material may occur over fibrous peat, etc. The following plant succession in the filling of a Minnesota lake has been suggested by Soper.<sup>1</sup>

1. Stonewort: waterweed stage.
2. Pondweed: water lily stage.
3. Rush: wild rice stage.
4. Bog: meadow stage.
5. Bog: heath stage.
6. Tamarack: spruce stage.
7. Pine association.

<sup>1</sup> "The Peat Deposits of Minnesota," by E. K. Soper, *Minnesota Geological Survey Bull.* 16, 1919.

# 2

## Processes of Soil Development

Having studied briefly the rocks and minerals which furnish the raw material from which soils are developed, we wonder about the processes which are active in converting this material into soil. As we would expect, these are largely the natural processes or forces in nature which bring about the breaking down, both physically and chemically, of objects all around us. We are familiar with the rotting of wood, the rusting of iron, the cracking of pipes or other containers as a result of the freezing of water, the solution of salts to give sea water its flavor, the formation of caves, etc. The processes which cause such phenomena are the same ones which, acting on soil material, result in soil development. However, there is a phase of soil formation other than the processes of destruction. These are the processes of construction, that is, the building up or addition of new materials. The action of all natural forces on soil material is included in the term *weathering*. The soil-forming processes may be considered under the following objectives.

### Objectives

- A. Processes of disintegration.
- B. Chemical processes of soil formation.
- C. Products and results of mineral decomposition.
- D. Biological activities and soil development.
- E. Constructive processes of soil development.
- F. General changes leading to soil development.
- G. The soil profile.

### PROCESSES OF DISINTEGRATION

Certain natural forces have a tendency to alter the shape and size of rocks and minerals without causing much if any change in the chemical composition of them. These are known as the physical forces of weather-

ing or disintegration processes. In arid regions the activities of disintegration processes are much more prominent than are the changes brought about by chemical processes because the lack of moisture retards chemical reactions. Disintegration processes may be considered in the light of two questions.

### Questions

1. How do temperature changes disrupt rocks?
2. In what ways does water aid in disintegration of soil material?

**Temperature Changes.** Different minerals usually have different coefficients of expansion: that is, their volume changes when heated and cooled are different. If, then, a rock composed of different minerals, like the granites, is heated and cooled repeatedly, it is literally pulled to pieces through the different volume changes of the minerals. In some arid regions where granitic rocks are plentiful, large accumulations of disintegrated granite develop, and the material is used on roads and for other purposes much as gravel is used in more humid climates.

Rocks are disrupted also through changes in temperature by having the outer shell expand more rapidly than the interior when warmed by the morning sun. During the day the rock becomes more or less warmed throughout and hence expanded. At night the outer portion contracts rapidly, owing to cooling after sunset, and hence the shell becomes too small for the interior. Repetition of these changes results in the cracking or scaling off of the rock exterior, as shown in Fig. 10.

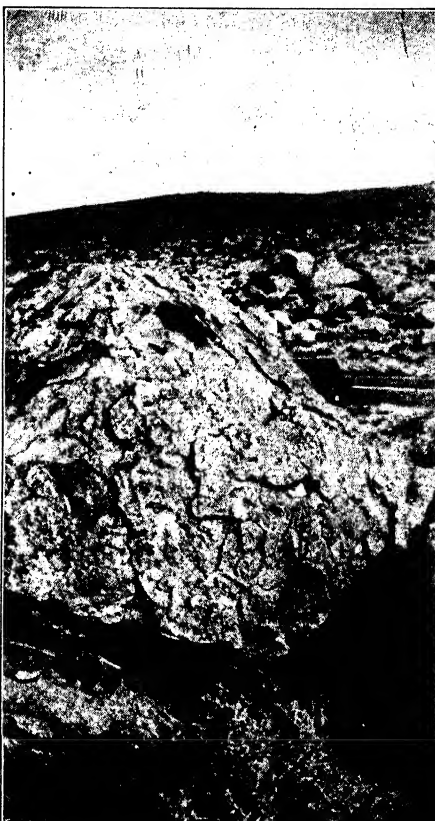


FIG. 10. Exfoliation or scaling off of the surface of rocks in the Mohave Desert due to changes in temperature.

**Water an Aid in Disintegration.** Upon freezing, water expands with a great force, approximately 150 tons to the square foot. Accordingly water, accompanied by changes in temperature, becomes an important agent in disintegrating soil material. As small cracks occur and crevices develop through solution or disintegration of compounds of calcium, iron, magnesium, and other basic elements, they are filled with moisture which alternately freezes and thaws, thus gradually enlarging the opening. Some rocks are so porous that they absorb sufficient quantities of water to shatter them upon repeated freezing.

Many substances also expand and contract appreciably upon wetting and drying. Thus serving as a wetting agent, water may contribute materially to the breaking up of soil material during various stages of soil formation.

### CHEMICAL PROCESSES OF SOIL FORMATION

Processes which change the chemical composition of soil material are termed the chemical forces of weathering. It should be remembered, however, that few changes in chemical composition occur without some change in the physical form of soil material. Accordingly, chemical and physical changes go hand in hand, but in this section our attention will be centered on the alterations in chemical composition.

In humid climates where the earth's crust is moist much of the time, chemical weathering is very active because moisture is necessary for chemical processes to proceed. Physical forces of disintegration are also operative, but the results of chemical changes overshadow their activities. The most active chemical changes are considered in answers to several questions.

#### Questions

1. How does oxygen function as a weathering agent?
2. What is meant by reduction?
3. What is the process of hydration?
4. How does hydrolysis function?
5. Is carbon dioxide an active agent?
6. In what way does solution aid soil formation?

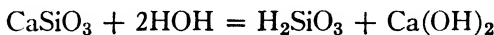
**Oxidation.** Oxygen is quite an active element, and it combines with many other elements freely. The uniting of oxygen with another element or an increase in the amount of oxygen in a compound is considered the process of oxidation in this discussion, although the chemist would consider this a very limited definition. Because oxygen makes

up about 20 per cent by volume of the atmosphere, there is abundant opportunity for oxidation of soil parent materials. This is one of the most common of the weathering processes. A simple example of oxidation is the union of carbon with oxygen to form carbon dioxide; thus  $C + O_2 = CO_2$ . A more complicated illustration showing how ferrous iron in a mineral may be oxidized to ferric iron is:  $4FeCO_3 + O_2 = 2Fe_2O_3 + 4CO_2$ . Here  $CO_2$  is split off.

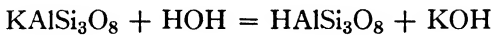
**Reduction.** As oxidation takes place in the presence of adequate oxygen, so reduction, the reverse of oxidation, occurs when the supply of oxygen is quite limited. This process, then, is active at lower depths in the earth's crust, under poorly drained conditions where water occupies much of the air space of the soil, on the interior of compact groups of very fine soil particles where air penetrates very slowly, and, in fact, wherever in the soil there is poor aeration. The evolution of  $CO_2$  from rapidly decaying organic matter may reduce oxygen supply and result in reduction.

**Hydration.** The union of water with soil minerals is illustrated by the *hydration* of iron oxide:  $2Fe_2O_3 + 3H_2O = 2Fe_2O_3 \cdot 3H_2O$ . Hydration is a common process in rock decay. Not only does the iron in minerals oxidize and become hydrated, thus increasing in size and softness, and so lead to further mineral breakdown, but also many other compounds, like  $SiO_2$  and  $Al_2O_3$  and aluminosilicates, as well as organic substances become hydrated.

**Hydrolysis.** Water is a very active chemical reagent. The H in water tends to change places with the positive ion (cation) in many compounds, thus forming two new compounds. This process goes on extensively with minerals containing strongly basic elements, parts of which are exchanged for H; thus,



and

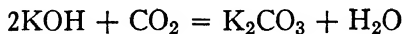


This exchange is known as *hydrolysis*, and it is a very effective reaction in changing the chemical composition of many minerals.

**Carbonation.** The decomposition of organic matter in the soil and the respiration of plant roots liberate large quantities of  $CO_2$  into the soil air. This gas readily combines with bases, producing carbonates or bicarbonates. For example, the  $Ca(OH)_2$  and  $KOH$  liberated by the hydrolysis reactions shown above readily combine with  $CO_2$  as follows:

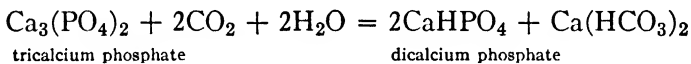
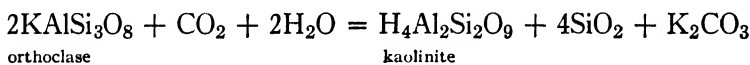


and



This process, known as *carbonation*, is one of the most effective in decomposing soil minerals. As the bicarbonates of calcium and magnesium are fairly soluble, carbonation is the primary process in the accumulation of residual soil material through decomposition of limestone and in the decomposition of minerals with a high calcium content.

The soil air contains approximately ten times as much CO<sub>2</sub> as atmospheric air. Furthermore, the film of moisture lying immediately next to the surface of the fine soil particles is highly charged with this gas. With this intimate association of oxygen, water, and carbon dioxide in contact with mineral particles, it cannot be expected that each will act independently on the soil minerals. On the other hand, oxidation, hydration, and carbonation usually proceed jointly or in such close association that the individual actions cannot be readily discerned. An example of the combined action of carbonation and hydrolysis is illustrated by the equations



**Solution.** The process of *solution* is also a chemical force of mineral decomposition, but, as pure water is a very weak solvent for most soil minerals, solution comes about largely through the joint action of  $\text{CO}_2$  and  $\text{H}_2\text{O}$ , or by water containing traces of mineral or organic acids, or by salt solutions. The principal mineral acids developed in the soil through chemical and biological agencies are  $\text{HNO}_2$ ,  $\text{HNO}_3$ ,  $\text{H}_2\text{SO}_4$ ,  $\text{H}_3\text{PO}_4$ , and  $\text{H}_2\text{SO}_3$ . Seldom do these acids occur in more than traces, but their continued formation ultimately results in considerable decomposition of minerals. Activities of soil organisms, such as bacteria and fungi, produce  $\text{CO}_2$  and complex organic compounds of acid nature. They also aid in the process of solution. Solutions of salts have a greater solvent action for many minerals than has pure water, and hence in soil areas where soluble salts are abundant the process of solution may be accelerated.

**PRODUCTS AND RESULTS OF MINERAL DECOMPOSITION**

The relative rate and predominance of the various processes of mineral disintegration and decomposition vary under different climatic conditions and as the actions proceed. In the early periods of the earth's history, when the surface was more or less continuous rock with little if any animal or plant life, the processes must have been mainly physical. As mineral fragments became smaller, exposing greater and greater surface to the action of water and oxygen and other gases which may have been present, chemical decay played a greater part. As animal and plant life developed, giving rise to  $\text{CO}_2$  and other decomposition products, chemical decomposition became increasingly active. In arid climates, however, where moisture is limited and where plant and animal life is not abundant, and in high altitudes where life is also limited, physical processes continue to predominate in mineral breakdown.

As soil material becomes more completely decomposed, all processes of physical and chemical decay become slower. How far these processes will continue is not determined, for no soil has yet been found which is not undergoing chemical and physical change. Soils that are approaching equilibrium with their environment or that are changing very slowly indeed are called mature soils.

It follows, then, that a logical concept of the nature of soils must necessarily take into consideration the dynamic characteristics of the materials from which, and in which, the soils are forming. A discussion of the results and products of the weathering processes described above serves as a helpful summary of these dynamic properties. In the discussion below the important points can be more easily distinguished if the following questions are kept in mind.

**Questions**

1. Is much material lost during soil formation?
2. What minerals and chemical compounds remain after rock decay?
3. How are soil properties affected by the processes and products of weathering?

**Losses during Weathering.** As pointed out in the earlier discussion, carbonate minerals are converted into bicarbonates, such as  $\text{Ca}(\text{HCO}_3)_2$  and  $\text{Mg}(\text{HCO}_3)_2$ , which are relatively soluble. These compounds may be removed from the soil if moisture and drainage conditions permit, or the cations may pass into other combinations and be retained. Fragments of the original minerals may remain for long

periods and make up an appreciable part of the soils subsequently formed. In other places the soils contain none of these fragments but are developed from the impurities that were in the minerals, such as sand, clay, and hydrated oxides of iron and aluminum. Thus, several hundred feet of limestone may be decomposed before enough residues are accumulated to make a soil a few feet in depth. The resulting soil may even be deficient in lime and hence acid in reaction.

The large accumulations of soluble salts in the ocean and saline lakes, the lime and other salts in hard water, our potash deposits and similar phenomena bear witness to the losses rocks undergo during decomposition.

**Minerals and Compounds that Remain.** Some minerals and rocks decompose readily, but others are quite resistant to weathering processes.

The oxide minerals remain largely in the soil as quartz and hydrated oxides of iron.

The micas and talcs remain as particles of various sizes, usually quite fine, and without great decomposition except when iron is a constituent.

The feldspars and other silicates decompose with the production of the carbonates and bicarbonates of calcium, magnesium, sodium, potassium, and other bases which vary in solubility. Some of the silica becomes soluble, and some remains as insoluble  $\text{SiO}_2$ . Much of the Si and Al remain as clay-like and sometimes jelly-like hydrated aluminosilicates as  $\text{H}_2\text{Al}_2\text{Si}_2\text{O}_8 \cdot \text{H}_2\text{O}$ . Hydrated oxides of iron and of aluminum are formed. With magnesium-bearing silicates, products, including hydrated magnesium silicates like  $\text{H}_2\text{Mg}_2\text{Si}_2\text{O}_8 \cdot \text{H}_2\text{O}$ , containing less Ca, Fe, and  $\text{SiO}_2$  than the original mineral are formed.

**Soil Properties as Affected by Products of Mineral Decay.** The processes of mineral decomposition result in the removal of the basic elements like Ca, K, Na, and Mg from the minerals and in the formation of more or less soluble compounds of them. A large part of the negative radicals of the minerals composed of silicon or aluminum and silicon remain as hydrated aluminosilicates, together with  $\text{SiO}_2$ ,  $\text{Al}_2\text{O}_3$ , and  $\text{Fe}_2\text{O}_3$ , in various stages of hydration. They are mixed with remaining particles of the minerals in different stages of decomposition. The physical effects of the process are to accumulate material in a much finer state of division, with part of it sticky or jelly-like because of hydration.

The chemical effects vary with climatic conditions. In areas of low rainfall the soluble products may accumulate near the surface, producing a soil rich in soluble salts and alkaline in reaction. In more humid

climates the soluble compounds containing the basic elements may leach away, leaving a soil poor in plant nutrients and in need of bases. Such soils are said to be acid in reaction. Under some climatic conditions the finely divided and hydrated oxides of iron and aluminum together with aluminosilicates may move downward, leaving the silica and undecomposed mineral particles in the surface layers. Under other climatic environments the silica and soluble salts may be carried downward, leaving an accumulation of hydrated iron and alumina.

### BIOLOGICAL ACTIVITIES AND SOIL DEVELOPMENT

The relative rate of activity of various soil-developing processes is greatly modified by the type of vegetation growing on the soil and by both the animal and plant life inhabiting it. At the same time the type of vegetation occupying a soil area and the animal life within it are influenced by the properties of the soil. For example, earthworms are scarce in certain soils of high acidity and low organic matter content. The work of fungi is said to predominate over bacterial activity in the coniferous forests of the cool, humid North. Certain species of plants grow poorly on acid soils, whereas others require high soil acidity for abundant growth. The interrelation of plant and animal life and soil properties is thus seen to be both intricate and interesting. A few of the most evident relationships between plants and animals and soil-forming processes are suggested in the following questions.

#### Questions

1. What forms of plants inhabited the earth's surface in the early stages of soil development?
2. Are bacteria, fungi, and similar life forms still active in soil formation?
3. How does the growth of higher plants affect soil properties?
4. Is animal life concerned in soil development?

**Activities of Elementary Plants.** In the very early stages of mineral decomposition and soil formation the lower forms of plant life, such as lichens, fungi, and bacteria, undoubtedly played a major role. The ability of lichens to grow on boulders and other rock surfaces exhibiting practically no indication of decomposition demonstrates their ability to extract mineral nutrients from unweathered minerals, to subsist with little water, and to obtain nitrogen from an undetermined source, possibly the air. Under clusters of lichens, even on very hard rocks, is found a thin accumulation of soil material, evidently the result of the decomposing action of the plant on the rock.

The ability of certain bacterial forms to extract nitrogen from the air for their own body needs, when supplied with mineral nutrients and carbonaceous food, and without associative growth of other plants, unquestionably played a considerable part in the building up of nitrogenous organic material in the soil in the early stages of development, thus paving the way for the growth of higher plants.

The work of these elemental plants has continued all during the process of mineral decomposition and soil development, and they remain active throughout the existence of the soil. It is through the action of such organisms that plant and animal remains are decayed, thus liberating the nutrients contained for the use of the next plant generation and simultaneously producing  $\text{CO}_2$  and complex organic compounds which aid in mineral decomposition.

**Influence of Higher Plants.** Higher plants, such as grass and trees, exert a great influence on the characteristics of the soil developed under them. Their influence on erosion and moisture relationships within the soil are contributing factors. The type of root development and the return of vegetable matter to the surface of the ground by both annuals and perennials play an active role in the nutrient cycle. The absorption of soluble nutrients from the soil and their return in plant parts, even the variation in chemical composition of leaves of different species, are important in their effects on soil development. Roots, particularly small ones, penetrate crevices in rock fragments, enlarging them and sometimes splitting the rock as growth continues. The porosity of the soil is also increased as the result of root growth. Some of these points are of sufficient significance to warrant a fuller discussion in later chapters.

**Work of Animals.** Worms, insects, and other small animal forms play a greater part in modifying chemical and physical properties of soils than do larger animals. The continuous burrowing of worms and insects establishes drainage channels in the soil and otherwise changes soil structure. Earthworms pass large quantities of soil through their bodies, thus altering it both chemically and physically. Worms and ants bring up a surprising amount of soil and deposit it on the surface. Many insects consume plant material both on the surface and within the soil. Worms are active in dragging fragments of leaves and grass into the soil to become a part of the soil organic matter. The abundant formation of "middens" in some timbered soils testifies to this activity of worms, whereas the disappearance from the soil surface of leaves of certain species indicates a strong choice on the part of worms as to the kinds of vegetation they consume.

**Additions to the Soil.** All types of animal and plant life living in the soil and soil material contribute to the soil substance. As time passes, the weight of plant roots buried in the soil is large, as is also the portion of the leaves and stems that is carried into and becomes a part of the soil. The bodies of animals, particularly insects and worms, contribute much organic substance to the soil material, and the weight of bacterial and fungi growth in an acre is unbelievably large.

### CONSTRUCTIVE PROCESSES OF SOIL DEVELOPMENT

Up to this point the discussion has been devoted primarily to the processes and products of disintegration and decomposition; in other words, to destructive processes. The conspicuous part of soil development is mainly the result of such processes, and they have received detailed study for many years. The constructive processes of soil development are not so evident to the eye and, therefore, have received much less study, although they may be more important in determining soil characteristics and the utilization of the soil for plant growth. Destructive and constructive processes, however, are intimately related. In fact, distinctions between the two are often slight. For example, a process may be destructive in its action on parent material and yet constructive in its effect on the resulting soil. There is, then, some confusion and inevitable overlapping in the use of these terms. This section emphasizes those processes that contribute most to the construction of soils even though they may have been discussed before as destructive to parent material.

Three questions are proposed as guides in studying the constructive processes of soil development.

#### Questions

1. What physical properties of soils are influenced by constructive soil processes?
2. What chemical compounds are developed in soil, and how do they affect soil properties?
3. What kind of organic compounds are built up in the soil?

**Soil Morphology.** The physical structure of soils presents more visible evidence of constructive action in soils than does chemical or mineralogical composition. The formation of zones of higher clay content, at various depths underlying more sandy or silty horizons, is a result of constructive change. The practical effects of such clay-rich horizons on moisture supply and root development are far-reaching. In soils of

sandy texture the additional moisture retained because of this horizon makes possible greatly increased yields of crops. On the other hand, if the clay layer is very dense, root development is restricted in places and the soil is too wet for the growth of some crops.

Other constructive soil-forming processes lead to the development of organic compounds that move downward and concentrate in definite zones. Hydrated oxides of iron are also constructive in their action. These substances sometimes cement the soil material into a semi-rock-like mass which limits root development and water movement.

Through freezing and thawing, the adhesive and cohesive action of moisture films, and the cementing power of sticky clay-like substances and salts, the mineral particles and fragments are bound together in aggregates of irregular but definite form and structure. Evidence of this structural arrangement is shown by the fact that earth removed from a hole will not completely fill the same hole when it is returned if plenty of water and tamping are employed. The development of a characteristic soil structure is a process requiring many years. From a practical standpoint structure is important because it influences moisture movements in both the plow layer and subsoil. The hard, massive layers and the clay horizons already mentioned are a part of structure. So are the channels formed by insects, worms, and plant roots. Not only do these and other structures control the speed of moisture movement, but they also control aeration. A farmer spends a large proportion of his time and labor in developing by means of tillage a soil structure favorable to root growth and rapid seed germination. Soil that loses its natural field structure loses its identity and becomes an entirely different medium for plant growth. The many different kinds of aggregates or structural units found in soil have been named and classified, and this knowledge constitutes a definite part of soil science.

**Chemical Compounds Developed in Soils.** As rocks and minerals decompose, portions of some of the products are recombined into new chemical compounds. There is a considerable list of *secondary* minerals which pedologists have so designated because they are formed through the decomposition of *primary* minerals. Clay is made up largely of such secondary minerals mixed with fragments of quartz ( $\text{SiO}_2$ ), hydrated oxides of iron, some organic matter, and other impurities.

The very fine clay, generally called colloidal material, is comprised principally of minerals developed in the soil. Three groups of these minerals have been recognized, namely, the montmorillonite group,  $(\text{OH})_4\text{Al}_4\text{Si}_8\text{O}_{20} \cdot \text{XH}_2\text{O}$ , the kaolinite group,  $(\text{OH})_8\text{Al}_4\text{Si}_4\text{O}_{10}$ , and the illite group,  $(\text{OH})_4\text{K}_x(\text{Al}_4 \cdot \text{Fe}_4 \cdot \text{Mg}_4\text{Mg}_6)(\text{Si}_{8-x} \cdot \text{Al}_x)\text{O}_{20}$ . In each

group there are several members which differ from the formula given primarily in the proportion of silicon and aluminum present. Also, there is considerable variation in composition of the members of the illite and montmorillonite groups owing to replacement of aluminum by ferric iron and sometimes magnesium. The crystals of these minerals are composed of plates of  $\text{SiO}_2$  and  $\text{Al}_2\text{O}_3$ , with varying degrees of hydration. The montmorillonite and the illite minerals have two plates of  $\text{SiO}_2$  and one plate of  $\text{Al}_2\text{O}_3$ , whereas the kaolinite minerals have one plate of each.

These minerals give the soil most of its cohesive and adhesive properties and are also very effective in holding certain plant nutrients in such a state that they are not readily lost by leaching and yet may become available for plant use. These materials are now receiving much intensive study and, as improved methods of procedure are developed and new scientific apparatus is perfected, much more information concerning these compounds, which are the result of constructive soil processes and which control many vital soil processes, will be gathered.

**Organic Compounds Formed in Soils.** The action of soil organisms in decomposing plant and mineral soil materials results in the development of many complex organic compounds which are active agents in soil processes. The bodies of the organisms themselves contribute appreciably to the organic content of the soil. These compounds not only serve as a means of storing a supply of nitrogen in the soil but also influence the chemical and physical soil properties. They affect soil structure, moisture retention and movement, and soil temperature to some degree. Furthermore, they function in retaining plant nutrients against leaching and in supplying them to plants.

## GENERAL CHANGES LEADING TO SOIL DEVELOPMENT

Many factors are active in the evolution of soils. Some of the processes are simple, many are complex, and all are more or less closely interrelated. Some changes proceed rapidly in the early stages of formation; others become dominant in the later stages of soil development. Most of the processes are overlapping, and all proceed simultaneously. The result is, of course, the soil. In reviewing the process of soil formation the following questions will be helpful.

### Questions

1. What changes in size of particles occur?
2. Do all minerals decompose at equal rates?

3. Does weathering attack the surface or interior of particles?
4. Which chemical elements are lost most rapidly from minerals?
5. Are new minerals developed in the soil?
6. What chemical changes occur in colloidal particles?
7. Is accumulation of organic matter a part of soil formation?
8. Is structure a soil characteristic?
9. How do environment and parent material influence soil characteristics?

**Main Steps in Soil Development.** The principal changes occurring during the development of a soil from or in its parent material may be summarized in the following statements:

1. The fragments or particles of parent material become smaller.
2. Easily decomposed minerals gradually disappear, leaving the more resistant minerals, the products of mineral decay, and secondary compounds.
3. As weathering attacks their surfaces, a coating of the residues of mineral decay accumulates around the particles. This coating consists of hydrated oxides of iron, aluminum, silicon, and hydrated aluminosilicates and frequently is jelly-like.
4. The basic elements are gradually removed from the surfaces of the particles and combined into compounds of variable solubility. In areas of high rainfall these salts are leached out, leaving the soil increasingly deficient in basic elements. In regions of less rainfall the salts accumulate in the surface soil or at various distances below the surface.
5. New minerals are developed which have definite crystal structures, although they occur in very minute particles, and which have a far-reaching influence on the chemical and physical properties of the soil.
6. In advanced stages of weathering the fine clay or colloidal particles attain a high hydrogen-ion content and a correspondingly low content of calcium, magnesium, sodium, and potassium ions. Some of the particles may then move downward until they are coagulated by different environmental conditions. This process gives rise to a zone or horizon of clay concentration.
7. Organic matter, in different stages of decay, accumulates in widely varying quantities on the surface and within the soil material.
8. A characteristic structure develops.

**Influence of Environment and Parent Material on Soil Characteristics.** Climate determines the relative rates of activity of the different forces of weathering and soil formation. It also limits the extent to which soil development proceeds. In arid regions, for example, the

lack of water has prevented soil development from progressing much beyond step 3. In these places where the moisture supply is limited, physical changes predominate. With increasing moisture content, chemical changes assume a more important role. Not only the moisture factor of climate but also temperature has an important influence on soil development. The speed of chemical reaction in humid regions varies markedly with the average seasonal temperatures and with the length of the warm season, as would be expected from the effect of temperature on the speed of chemical reactions. Climate also plays a major role in determining the distribution of different kinds of vegetation.

In the early stages of soil development the nature of the parent material dominates the characteristics of the soil. Such soils are said to be young. But, when parent material has been subjected to soil-forming processes for long periods of time, the soils of similar climatic zones develop some similarities independent of the kind of parent material. These soils that reflect the characteristics commonly indicative of the climate and associated vegetation are called zonal soils.

Since weathering and soil-forming processes start at the earth's surface and proceed downward, the results of these processes give rise to layers more or less parallel to the land surface. These layers are called *horizons*.

### THE SOIL PROFILE

As previously stated, the result of soil-forming forces working for long periods of time is the gradual differentiation of layers or horizons within the parent material. Collectively these horizons are called a *profile*. The following questions emphasize the important points in studying soil profiles.

#### Questions

1. How are soil horizons named and described?
2. How does topography influence the profile that develops in a given parent material?
3. How does drainage influence soil-profile development?
4. What is the relation between soil reaction and soil development?
5. Is the composition of the parent material important?
6. How do the *A<sub>2</sub>* and *B* horizons vary in different soils?
7. Does organic matter affect profile characteristics?
8. Why are salt accumulations found in some soils?
9. What is a soil series?

**Differences in Horizons.** There are many kinds of soil profiles because no two horizons are ever exactly alike. Not only does their mineral composition vary, but also the percentage of different-sized particles as well as their structure and color differs widely within short distances. The nature of the horizons in a profile and the intensity of their development depend on many factors, a few of which are the composition of the parent material, length of time it has been exposed to soil-forming processes, climate, nature of the vegetative cover, and site characteristics such as relief and topography. Most profiles, however, are generally characterized by two or more of the following horizons: (1) The surface which consists of or contains an accumulation of plant leaves, stems, and roots; this horizon of plant residues varies from less than an inch to 20 inches or more in thickness; (2) a horizon which, because of the action of soil-forming forces, has lost part of its original constituents; (3) a horizon of accumulation (this is the layer where material removed from the two overlying horizons is deposited); (4) the horizon of parent material the upper portion of which may have undergone a small amount of alteration (as a whole, this material is assumed to be similar to that which existed on the surface before soil-developing processes began).

In pedology these horizons are conventionally designated by letters. The surface layer, or *A* horizon, is subdivided into the *A<sub>00</sub>*, *A<sub>0</sub>*, *A<sub>1</sub>*, *A<sub>2</sub>*, and *A<sub>3</sub>* sections.

The *A<sub>00</sub>* refers to the undecomposed litter of twigs and plant leaves; in the *A<sub>0</sub>* only partially decomposed organic matter is found. That layer where mineral particles predominate but the dark color of organic residues persists is designated as the *A<sub>1</sub>* horizon. The *A<sub>2</sub>* layer, on the contrary, is comparatively light in color and shows the maximum effects of leaching or *eluviation*.

The horizon of accumulation or *illuviation* is technically designated as the *B* horizon. It is often subdivided into the *B<sub>1</sub>*, *B<sub>2</sub>*, and *B<sub>3</sub>* sections, depending on the degree of accumulation in evidence. The parent material is referred to as the *C* horizon, and underlying strata (such as bed rock) are called the *D* horizon. These terms and their relationships are presented in Fig. 11. Notice that horizons *A* and *B* together constitute the *solum*.

As previously stated, all these horizons are not present in every soil. Fig. 12b shows a profile in which the light-colored *A<sub>2</sub>* horizon is not evident. Insufficient rainfall and a grass vegetation are two factors which contribute to the development of this type of profile.

In some cases neither the *A<sub>2</sub>* nor the *B* horizon is discernible, and a

profile similar to the one in Fig. 12c is produced. A high water table which has limited the activity of weathering agencies, a limited rainfall,

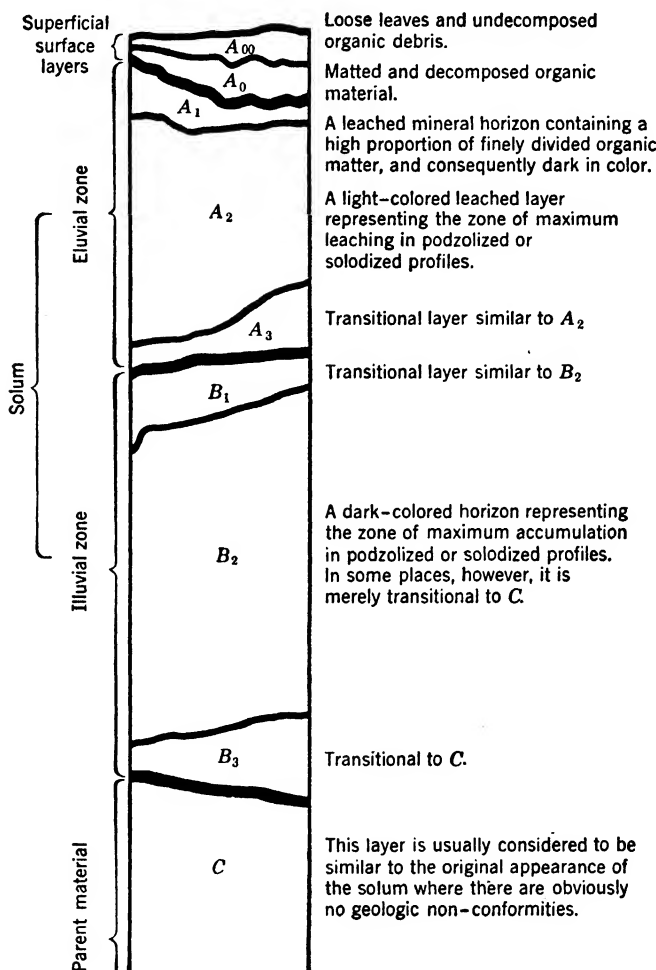


FIG. 11. A generalized profile of a timbered soil developed in a humid climate with moderate temperature. The thickness of the various horizons varies as indicated.

and a comparatively short period of activity of soil-developing processes are some of the conditions that give rise to this type of profile.

**Influence of Topography on the Soil Profile.** Topography modifies the soil-profile development in three ways: (1) by influencing the

quantity of precipitation absorbed and retained in the soil, thus affecting moisture relations; (2) by influencing the rate of removal of the soil by erosion; and (3) by directing movement of materials in suspension or solution from one area to another. As moisture is essential for the action of the chemical and biological processes of weathering and effectively acts in conjunction with some of the physical forces, it is

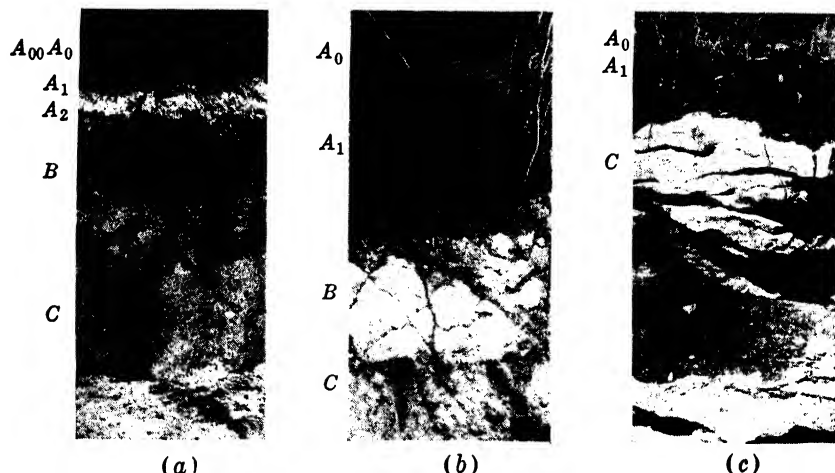


FIG. 12. Compare the thin  $A_1$  and the strongly developed  $A_2$  horizons of the Podzol profile on the left with the deep  $A_1$  and the absence of  $A_2$  horizon in the Chernozem profile in the center. Profile *c* shows the profile of a soil developing from limestone. Note the absence of a *B* horizon. [Profiles *b* and *c* by courtesy of Dr. Harper of the Oklahoma Experiment Station.]

evident that a modification of moisture relationships within a soil will materially influence profile development. In a humid glaciated region it is noteworthy that, in similar parent material of intermediate or fine texture, the soil of steep ridges or hills differs from the soil on gentle slopes or on a level to undulating topography. In arid climates these soil differences associated with differences in slope are much less pronounced.

On steep slopes the continuous removal of surface soil by erosion keeps exposing the lower horizons and so modifies the profile. Furthermore, the decreased thickness of the humus-containing soil results in a lessened supply of the products of decay of organic matter which contribute to the decomposition of minerals and to horizon-forming processes. Consequently the soils on steep slopes have thinner solums and

less conspicuous horizons than soils on level or undulating topography when the water table is well below the solum. These profile differences due to slope are least pronounced in soils developed in coarse-textured parent material in which internal drainage is excessive.

Topography indirectly plays another part in profile development by influencing the supply of moisture available for plant growth. It also

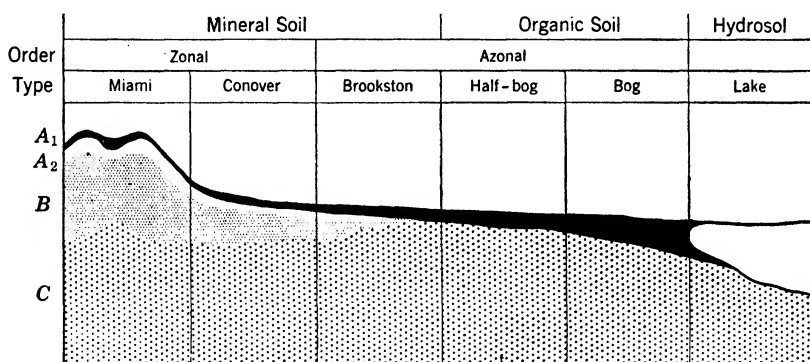


FIG. 13. Topography, through its influence on drainage, is a potent factor in soil development in humid regions. The moderate humus accumulation, well-defined horizon of eluviation, and strongly developed *B* horizon of the zonal profile are superseded by an increase in humus, diminution in eluviation, and accumulation in the *B* horizon as poorer drainage limits the action of soil development processes. Finally the bog is encountered, and then the hydrosol or body of water itself. In the diagram the thickness of the *A*<sub>1</sub> and *A*<sub>2</sub> horizons is intentionally exaggerated to illustrate the effect of topography and drainage on them.

has a bearing on the agricultural value of the land because it is related not only to both external and internal drainage conditions but also to the ease of performing tillage operations.

**Effect of Drainage Conditions on Soil Development.** As pointed out before, drainage materially influences soil-forming processes. Poor drainage may have even more effect on soil development than excessive drainage. The accumulation of organic matter is usually facilitated because it is preserved by water. Also, because of their low topographic position, poorly drained soils generally receive both organic and mineral matter that has eroded from the adjacent slopes. In arid regions soluble salts also accumulate in areas receiving drainage water from surrounding soils. Colors of the soil material at shallow depths are changed from yellows, reds, and browns, denoting good aeration

and oxidizing conditions, to the drabs, grays, and mottled yellows resulting from chemical reduction where drainage is poor. The horizon of eluviation is modified or may not be evident at all because of the slow or infrequent downward movement of water, and the *B* horizon is often replaced by a gray or bluish gray horizon known as *gley*. In this layer iron compounds as well as compounds of calcium, magnesium, and manganese are changed to soluble forms. In some places, however, the underlying layers may be so slightly changed that the surface horizon appears to rest directly on the parent material.

Intermediate drainage conditions give rise to other profiles. A water table, for instance, at a depth of 2 to 3 feet below a nearly level surface in humid areas, encourages the development of a conspicuous eluviated horizon underlain by an intensely developed illuviated layer. At times this horizon is indurated or cemented into a hard rock-like mass. In arid areas where ground waters frequently contain salts, alkali soils may result under these drainage conditions.

As a result of differences in topography or drainage or both, the soil profiles developed in similar parent material of a like age and within a single zonal region vary appreciably. A group of soils developed under such conditions and showing such variations in profile characteristics is designated as a *catena* or *toposequence of soils*.

**Soil Reaction a Characteristic of Soil Profiles.** Parent material ranges from that composed largely of acid minerals, through that containing various percentages of limestone, to limestone itself. Compounds containing calcium and magnesium, which give it an alkaline reaction, make up varying proportions of the parent material. As the processes of chemical weathering proceed, these basic elements are transformed into soluble compounds that either accumulate in lower horizons under arid conditions or leach out to a greater or lesser extent in more humid climates. In time, then, either the soil horizons may become more basic in reaction or they may become acid, depending on the prevailing climatic environment. Accordingly, within a single region a soil-reaction test may give some indication of the extent to which soil development has progressed.

The reaction of a soil horizon also indicates chemical conditions which are conducive either to the flocculation of the finer clay, on the one hand, or to its dispersion, with the possibility of being carried downward by water, on the other. The formation of soluble compounds of iron, aluminum, silicon, manganese, and many other elements is also related to soil reaction. Moreover, the activity of bacteria and other

soil organisms as well as the growth of higher plants is affected by soil reaction. A determination of the reaction of the various soil horizons is therefore useful in studying profile development and soil properties.

**Composition of Parent Material Important.** The composition of the parent material has much to do with the characteristics of the

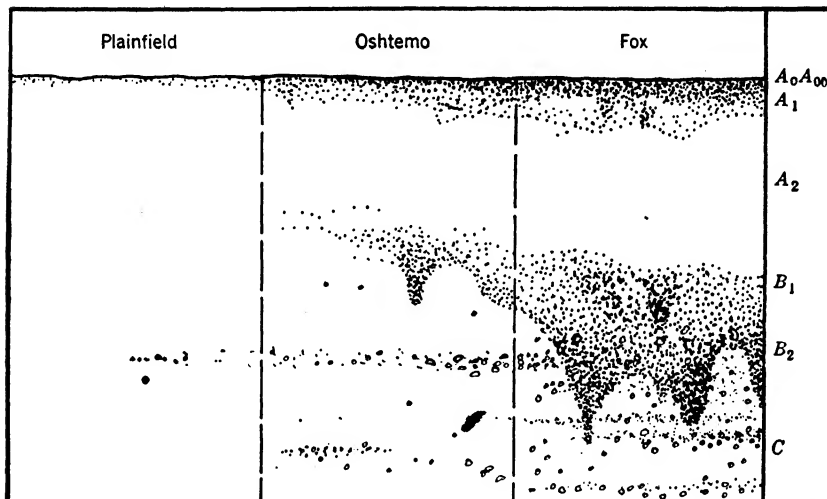


FIG. 14. The chemical composition of coarse material, deposited as an outwash plain, greatly influences the characteristics of the soil subsequently developed. The material on the left was highly siliceous and resulted in a soil having a minimum of clay in the *B* horizon. The Oshtemo soil developed a well-defined *B* horizon containing some clay because of the presence of appreciable quantities of limestone fragments in the parent material. The outwash on the right contains considerable quantities of limestone, and hence the Fox soil has a strongly developed *B* horizon with sufficient clay to bind the sand and gravel into a water-retentive layer which gives the soil a much greater agricultural value than is possessed by the Oshtemo or Plainfield.

profile developed, at least until the soil becomes very old indeed. If the material contains a large proportion of minerals which decompose with relative ease, there will be much clay produced, some of which, under suitable conditions, will accumulate in the *B* horizon, thus making a heavier "subsoil." On the other hand, if the soil material is composed almost entirely of minerals which weather very slowly, there will be very little accumulation in the *B* or illuviated horizon. In nature we find all variations between these extremes. Figure 14 illustrates the

effect of difference in composition of outwash plain sediments on the nature of the soil profiles developed.

**Variation in Properties of the Lower Soil Horizons.** The possibilities of variation in the horizon of illuviation have already been mentioned. In some sandy soils it contains very little more clay than the parent material and differs from adjoining horizons mainly in color. This kind of a *B* horizon contributes little to the crop-producing capacity of the soil. In other sandy soils, however, the illuviated horizon is higher in clay than the other horizons and hence serves as a reservoir for the storage of moisture for crop use. In some cases the *B* horizon may be either so firmly cemented or composed of such a dense accumulation of clay that plant root growth and water storage are restricted to the *A* horizon. On the other hand, in many soils derived from heavy clay the *B* horizon has a more open structure and is more permeable to roots than the parent material. Accumulations of calcium and other nutrients and concentrations of humus sometimes are found in the horizon of illuviation. Of course, these substances add materially to the fertility of the soil when present in correct amounts in the *A* horizon. However, when concentrated in the *B* horizon they may act like the clay accumulations mentioned above. The hardpan soils of arid regions are examples of soils with calcium-cemented subsoils, and the ground water podzols of humid areas have cemented subsoils high in organic matter and iron oxides.

The thickness of the *B* horizon is also an important property of a soil. A thin layer has little influence, favorable or unfavorable, on plant growth, but a thick layer of the same nature may decidedly affect the crop-producing power of the soil.

The *A<sub>2</sub>* (eluviated) horizon is frequently lower in clay content, more acid, and poorer in nutrients than the other horizons of the profile. The thickness of this horizon, therefore, influences the fertility of the soil. This horizon is usually absent in soils of arid regions. In many soils the light-colored layer is not a good medium for rapid seed germination or for the development of young roots. As the *A<sub>1</sub>* horizon, or dark-colored plow layer decreases in thickness through erosion and more and more of the underlying eluviated horizon is incorporated in the plow layer, it therefore becomes necessary to supply additional quantities of organic matter, manure, and commercial fertilizers to maintain the productivity of the land.

**Organic Matter a Distinguishing Feature of Soils.** The quantity of organic matter that accumulates in soil depends on many factors, such as the length of growing season, soil-moisture relationships, type

of vegetative cover, soil reaction, temperature, rainfall, and nutrient content. Because all these factors affect other soil characteristics, the amount of organic matter is often considered an index of soil conditions. Organic matter, through both its presence and its decomposition products, exerts so great an influence on soil properties that the quantity becomes an important consideration in comparing soil profiles.

**Salt Accumulation in Soils.** In climatic regions where precipitation percolating through the decaying minerals is not sufficient to carry away the soluble compounds developed or present in the parent materials, they accumulate as salts in the profile. The depth and extent of these accumulations together with their chemical composition play important parts in determining soil properties and soil utilization. Accumulations of salts in soils occur chiefly, but are not confined to, arid climates; occasionally they are also found in soils of humid regions.

**Soil Series.** It has been found that in spite of the differences that exist in soil profiles they are frequently very similar over considerable areas of the earth's surface. Closely similar profiles are grouped together as a *series*. The profiles of a series are similar in origin; that is, they developed under the same conditions of climate and vegetative cover, through the same forces acting on similar parent material for approximately the same length of time. Topographic and drainage conditions associated with the profiles in any particular series are similar. The corresponding horizons in the different profiles within the soil series are similar in depth, structure, color, reaction, and chemical characteristics. The organic matter content as well as the nature and supply of humus is also similar.

Variations within the series are confined to difference in texture of the surface horizon or plow soil. Thus there may be loams, silt loams, and clay loams within a series. At present, however, the tendency is to restrict this grouping to closely related textural classes. Clay soils and sandy soils do not, therefore, occur in the same series; few are even broad enough to include loams, sandy loams, and loamy sands.

A soil series is usually given the name of a town, river, or other geographical or political feature near which it was first identified. The Miami series, for example, was first recognized near Miami, Ohio.

# 3

## Physical Properties of Soils

The physical properties of a soil have much to do with its suitability for the many uses to which man puts it. The rigidity and supporting power, both wet and dry, the freedom of drainage, moisture-storage capacity, plasticity, ease of penetration by roots, aeration, and retention of plant nutrients are all intimately connected with the physical condition of the soil. It is pertinent, therefore, that persons dealing with soil in any way should be acquainted with the physical nature of different soils, knowing to what extent and by what means man can alter those properties—whether it be as a medium for plant growth or as a structural material in the making of highways, dams, and foundations for buildings or other engineering purposes, in manufacturing brick and tile, or in building golf courses, athletic fields, and running tracks.

The study of the physical properties of soils will be presented under eight topical divisions.

### Objectives

- A. Making a mechanical analysis.
- B. Properties of soil separates.
- C. A textural grouping of surface soils.
- D. Soil structure.
- E. Tillage operations and soil properties.
- F. Porosity and weight of soil.
- G. Soil color.
- H. Soil temperature.

### MAKING A MECHANICAL ANALYSIS

The physical properties of soils are dependent mainly on the surface exposed by the particles and on the content of organic matter, although the chemical properties of the particles are also a contributing factor. The surface exposed by the particles is primarily a function of their size,

and hence the study of the physical properties of a soil may well begin by determining the percentage of the different size groups or separates present; in other words, by making a mechanical analysis. The significant points relative to the making of mechanical analyses of soils are indicated by the following questions.

### Questions

1. What are soil separates?
2. How can the tendency for the particles to stick together be overcome?
3. How are the sands separated from the silt and clay?
4. By what means are the different size groups of sand separated?
5. What is the procedure for separating silt from clay?
6. What is the hydrometer method?
7. What is the procedure for the pipette method?

**Soil Separates.** One of the first facts to attract the attention in examining soils is that they are composed of particles of different sizes. Needless to say, the proportions of different-sized particles in soils have much to do with their physical and chemical properties, and hence with plant relationships.

Soil particles have been divided into groups entirely on the basis of size, that is, without regard to their chemical composition, color, weight, or other properties. These groups of particles are termed soil *separates* because any soil composed mainly of mineral matter may be *separated* into these particle groups. The process of dividing a soil into its separates and determining the relative proportion of each separate present is called *mechanical analysis*. In Table 1 are given the names of the separates, together with their diameters, the number of particles in 1

TABLE 1  
SOME CHARACTERISTICS OF SOIL SEPARATES

| Name           | Diameter,*<br>millimeters | Number<br>of Particles<br>per Gram | Surface Area in<br>1 Gram,<br>square<br>centimeters |
|----------------|---------------------------|------------------------------------|---|
| Fine gravel    | 2.00-1.00                 | 90                                 | 11.3  |
| Coarse sand    | 1.00-0.50                 | 722                                | 22.7  |
| Medium sand    | 0.50-0.25                 | 5,777                              | 45.4  |
| Fine sand      | 0.25-0.10                 | 46,213                             | 90.7  |
| Very fine sand | 0.10-0.05                 | 722,074                            | 226.9   |
| Silt           | 0.05-0.002                | 5,776,674                          | 453.7   |
| Clay           | below 0.002               | 90,260,853,860                     | 11,342.5  |

\* As established by U. S. Department of Agriculture.

gram (they are assumed to be spheres with maximum diameter of the group), and the surface area in square centimeters exposed by the particles in 1 gram of the separate.

**Dispersing the Soil.** Before a soil can be divided into groups of particles on the basis of size, it is essential to overcome the tendency of the very small particles to cling to the larger ones and to each other. Every particle must exist as an individual. Organic matter is one of the chief agents which bind particles together, and hence the first problem is to destroy this cementing material. The commonly accepted procedure is to oxidize the organic matter by boiling the sample in hydrogen peroxide solution.

Other cementing agents are calcium carbonate and the oxides of iron and aluminum. The carbonate may be dissolved by treatment with dilute hydrochloric acid. Iron and aluminum oxides also may be dissolved with suitable chemical reagents, but it is doubtful if such a procedure is advisable as the oxides are a part of the very fine or colloidal clay and should be determined as such.

The usual procedure is to shake the sample, after removal of the organic matter, in water in order to surround each small particle with a film of water which helps keep the particles apart. Usually a chemical such as sodium oxalate is added to the suspension in order to replace calcium, magnesium, and similar elements having a tendency to cause the fine particles to gather in clusters (flocs). The sodium combines with the fine clay particles and decreases their tendency to cohere. The sample may be shaken moderately for several hours, or it may be stirred violently for a few minutes in a stirring machine similar to those used in making malted milk drinks.

**Separating Sand from Silt and Clay.** The sand particles may be separated from the silt and clay by shaking the dispersed sample in a tall cylinder of water and decanting or siphoning off the turbid liquid when the sands have settled. Several decantations are necessary to effect complete separation.

**Separation of Sands.** The simplest procedure for separating the different-sized sand groups is to use carefully calibrated screens.

**Silt and Clay Separations.** Usually clay is separated from the silt by the principle of sedimentation, but settling is generally facilitated by means of centrifugal force. The suspension of silt and clay is placed in the tube of a special centrifuge which is run for a given time at a speed sufficient to throw the silt to the bottom of the tube, leaving the clay in suspension.

**The Hydrometer Method.** Bouyoucos has devised a method of determining the silt, clay, and sand content of a soil without separating them. The sample is dispersed in a stirring machine making use of a dispersing agent. The suspension is then poured into a tall cylinder, and a specially designed hydrometer is placed in it. By reading the hydrom-

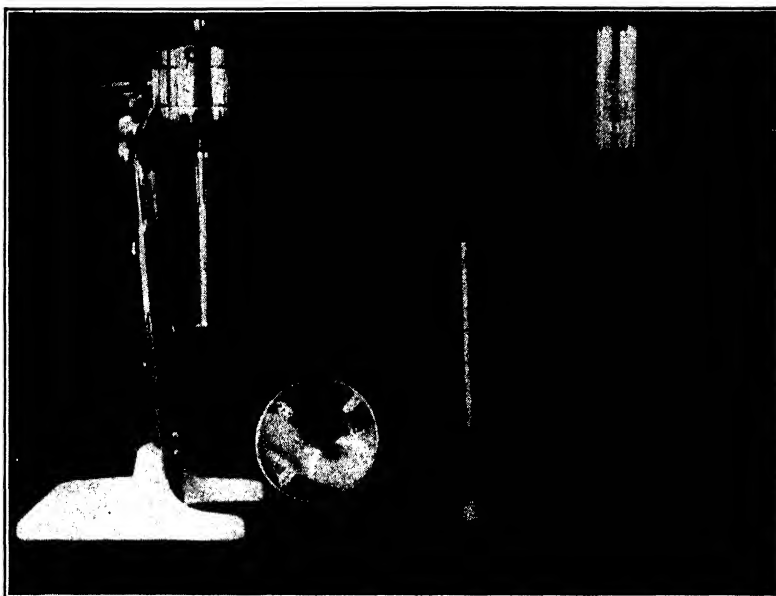


FIG. 15. Equipment used in making a mechanical analysis by the hydrometer or Bouyoucos method. The cup in which the soil is stirred is inclined to show the baffles which are essential if complete dispersion is to be obtained.

eter at the time interval (40 seconds) required for the sand (greater than 0.05 mm. in diameter) to settle, the quantity of silt plus clay in suspension can be determined, and hence the quantity of sand by difference. When sufficient time (2 hours) for the silt (greater than 0.002 mm. in diameter) to settle has elapsed, the percentage of clay may be determined by reading the hydrometer, and so the quantity of silt calculated. The very fine or colloidal clay is determined after a 24-hour period of sedimentation by the use of an especially sensitive hydrometer.

**The Pipette Method.** In this method the dispersed suspension is poured into a standard-sized cylinder, and after a predetermined time a given volume of the suspension is drawn off from a given depth by means of a pipette. The withdrawn portion is evaporated and the sedi-

ment weighed. Thus the density of the suspension is determined at a given depth as a function of time. By repeating these measurements at given time intervals the percentage of particles present with different settling velocities can be determined. As the settling velocity varies with particle size, the percentage of particles of different sizes in the sample may be determined by calculation.

Other methods for making a mechanical analysis of a soil have been suggested. The methods described, however, are those in most general use.

### PROPERTIES OF SOIL SEPARATES

Soils are divided into textural classes on the basis of the proportion of the different separates they contain. To understand the properties of the soil classes, it is essential to have a knowledge of the properties of the different soil separates. Except for the colloidal fraction the interiors of the soil particles are not exposed, and hence their chemical and physical activities must take place on the surface of the particles. The surface exposed by particles becomes, accordingly, of primary importance in a study of their properties. The very fine particles, because of their crystal structure, have an active internal as well as external surface. The questions below are submitted as guides in the study of the soil separates.

#### Questions

1. What are the chief functions of sand in soils?
2. Is silt an active soil constituent?
3. In what ways does colloidal clay differ from other clay?
4. How are the clay minerals constituted?

**Functions of Sand.** Sand particles are of comparatively large size and hence expose little surface compared to that exposed by an equal weight of silt or clay particles. Because of the small surface of the sand separates, the part they take in the chemical and physical activities of a soil is almost negligible unless some of the particles are composed of  $\text{CaCO}_3$ ,  $\text{MgCO}_3$ , or some other appreciably soluble compound. The smaller sand particles may contain a sufficient coating of the very small clay separate to attain some activity, but the action is due really to the clay and not to the sand. Because the sands are inactive, their chief function in soil is to serve as a framework around which the active part of the soil is associated.

Unless present in too small proportion the sands increase the size of spaces between particles, thus facilitating movements of air and

drainage water. They also add considerably to the weight of the soil, as will be pointed out later in the discussion of soil structure and volume weight.

**Silt in Relation to Soil Properties.** The coarser silt particles are very similar to the finer sands in surface exposed and hence take very little part in the chemical activities of soils. The finer silt has sufficient surface to give it some slight chemical activity and, if present in considerable quantity, must be considered in the determination of the total activity of the soil. There is an appreciable tendency for the very fine clay to adhere to the surface of the silt particles which derive some activity from this source.

The minerals composing the silt particles have undergone little decomposition in certain soils, primarily those in areas of limited rainfall and in soil derived from recently deposited glacial or loessial material. In such soils the silt may yield appreciable quantities of nutrients to the soil solution or to the colloidal clay.

In themselves, silt particles have little tendency to stick together or to adhere to other particles and hence have little influence on the particle arrangement or structure of the soil.

**Clay vs. Colloidal Clay.** The term clay is rather loosely used. In early methods of mechanical analysis all particles with a diameter less than 0.005 mm. were considered clay. Further investigation showed that the largest of these particles, those with a diameter of 0.002 mm. and greater, had comparatively little activity, and hence it was argued that they should be included in the silt separate. A given weight of particles with a diameter of 0.0001 mm. has 20 times the surface of particles with a diameter of 0.002 mm. and, accordingly, is very much more reactive. On the basis of activity, it has seemed advisable to divide the clay fraction into two groups, calling the finer material colloidal clay (from the Greek words *kolla*, meaning glue, and *oid*, meaning like) and setting 0.0002 mm. as the upper limit of diameter for this material. This size limit is the subject of considerable discussion and chemists are inclined to place it lower. The laboratory for chemical and physical investigation of soils, U. S. Department of Agriculture, uses 1 micron<sup>1</sup> as the upper limit.

The coarser clay, that ranging from 2 microns to 200 millimicrons in diameter, has considerable reactivity and plays a definite part in the

<sup>1</sup> 0.001 mm. constitutes 1 micron or  $\mu$ ; and 0.000001 mm., 1 millimicron (1 m $\mu$ ). 200 m $\mu$  would then be equal to 0.0002 mm. For convenience these designations of size will be used in referring to clay and colloidal clay.

physical and chemical properties of soil. It is particularly effective when acting in association with the colloidal fraction.

**Clay Minerals.** The sand and silt of a soil are composed primarily of fragments of dense or non-porous minerals. Clay, on the other hand, and especially colloidal clay, is made up largely of minerals built up or synthesized in the soil. A chemical analysis of clay shows it consists largely of silicon, aluminum, and water, with varying quantities of iron, calcium, magnesium, potassium, and sodium. Small amounts of other elements may be present. The minerals are built up of layers or plates of  $\text{SiO}_2$  and  $\text{Al}_2\text{O}_3\text{XH}_2\text{O}$ . Sometimes portions of the Al are replaced by Fe or Mg. The kaolinite group<sup>2</sup> of minerals consists of one layer of silica and one of alumina, whereas the montmorillonite group is built of two plates of silica and one of alumina. The constitution of these minerals is discussed more fully on pages 107-108. X-ray diffraction shows these minerals to have definite crystal structure with a large amount of space between atoms composing the crystal. The crystal composed of three plates has much more internal space than that made up of two plates. The internal surface of these crystals added to the exterior surface gives them a much greater activity than they would otherwise possess.

The shape of the clay minerals is also a contributing factor to their activity. If they are plate-like in structure, as they usually are, they have much more surface than if they are spherical or cubical. The shape of these minerals also determines the manner in which they pack together. Plate-like forms adhere tightly when they are arranged flat side to flat side.

#### A TEXTURAL GROUPING OF SURFACE SOILS

The casual observer is inclined to confine his observations of soil properties to the surface soil or that portion turned by the plow, paying particular attention to color and texture. Texture refers to the proportion of different-sized particle groups in the soil on a percentage basis. This property is important because it is directly related to the total

<sup>2</sup> The minerals of the kaolinite group in addition to kaolinite are dickite, halloysite, metahalloysite, and nacrite. These have a 1:1 crystal lattice (one plate of  $\text{SiO}_2$  to one plate of  $\text{Al}_2\text{O}_3$ ) and are fixed or non-expanding. The montmorillonite group includes also beidellite and nontronite. They have a 2:1 crystal lattice that expands on wetting. A third group of clay minerals with a 2:1 fixed crystal lattice is known as the illite group. Specific names have not yet been given to the members.

surface of the soil particles and therefore to many phenomena in soils that have a bearing on management practices and productivity.

The steps followed in developing a grouping of soils based on particle size are indicated by questions.

### Questions

1. What are soil classes?
2. What are the names and distinguishing characteristics of several common soil classes?
3. Is there a generalized textural grouping of surface soils?
4. Does the class name convey any information concerning a soil beyond the percentage of separates contained?

**Soil Classes.** Never is a soil composed of only one separate. Usually at least small quantities of the majority of the separates are present. The next step in a textural classification, therefore, is to group them on the basis of the proportion of the different separates present. These groups are designated as soil *classes*, which are named according to the separate or separates which contribute most to their characteristics. This does not mean that a class is necessarily named after the separate present in largest quantity. It takes a very large quantity of coarse particles to exert as much influence on soil properties as a comparatively small quantity of the finest particles—clay. Clay is the most potent separate in imparting its properties to a mixture of separates, and hence the adjective clay is found in the class name of many soils which contain a higher percentage of the other separates than they do of clay.

It is customary to describe a soil horizon by the class name which is in accord with the physical make-up of that layer of soil. In the name of a soil type only the texture of the surface or plow soil is designated. But there is frequently a rather close relationship between the texture of the surface soil and that of the soil parent material. For example, a sandy surface soil frequently rests on material high in sand content. In youthful soils the texture of the whole soil profile may be similar to that of the parent material. However, there are so many exceptions to these relationships between the surface texture and the characteristics of the underlying soil horizons that these are very dangerous generalizations, as many people have learned by bitter experience.

**Common Soil Classes.** Figure 16 gives the proportions of the separates in some of the classes commonly used in describing soils. The sum of the percentages of sand, silt, and clay at any point in this diagram

## Physical Properties of Soils

is 100. For example, point *A* represents 15% clay, 65% sand, and 20% silt, and the textural or class name of this sample is sandy loam.

A soil composed of a mixture of separates such that the properties of no one separate or group of separates dominates its characteristics is called a *loam*. The stickiness of the clay and the floury nature of the

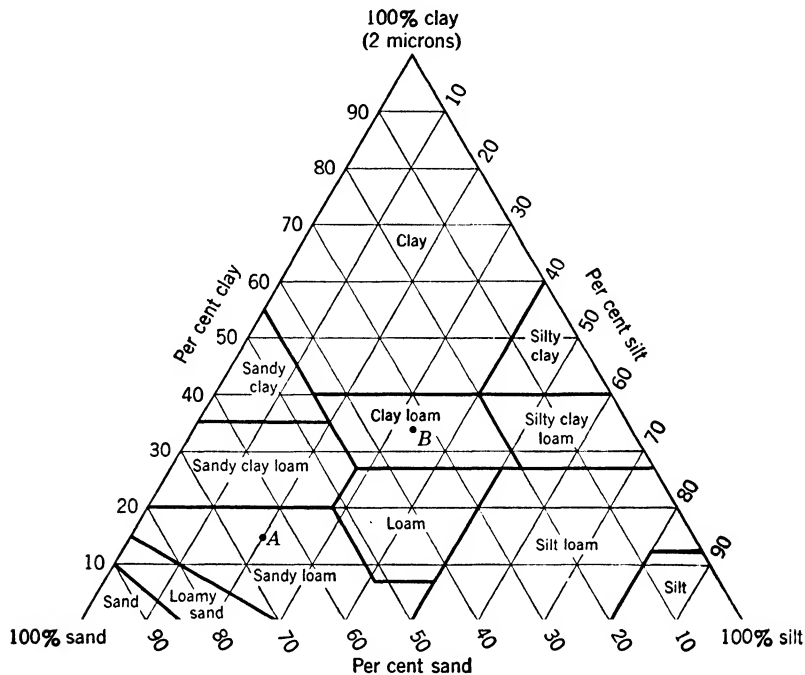


FIG. 16. The percentages of sand, silt, and clay in the textural classes of soil may be determined from this diagram. [Courtesy of Bureau of Plant Industry, Soils, and Agricultural Engineering, U.S.D.A.]

silt are balanced by the non-sticky and mealy or gritty characteristics of the fine to coarse sands. Water drains through the soil with ample freedom, and yet considerable moisture is retained for plant use. A loam can easily be kept in a state of good tilth which is conducive to a rapid germination of seeds and to easy penetration by roots.

Where a soil of loam-like qualities contains sufficient sand separates so that their characteristics predominate over those of silt and clay, the texture is a *sandy loam*. If the quantity of one of the sand separates is adequate to impart the properties of that separate to the soil, the name of the separate is added to make the soil class name. Thus,

in a *fine sandy loam* the qualities of fine sand predominate over those of the other sand separates; in a *sandy loam* the properties of the medium sand are evident. If a soil contains very little more silt and clay than are necessary to throw it into the sandy loam group, it is frequently called a *light sandy loam*. On the other hand, in a *heavy sandy loam* the sand content approaches the lower limit, and the silt and clay content the upper limit for a sandy loam.

Soils with a silt and clay content just below that required for sandy loam are generally designated *loamy sands*. This name was adopted because such soils produce moderately good yields of crops when rainfall is adequate, and therefore it seemed undesirable to classify them as sands which are decidedly lower in productivity characteristics.

Soils containing even less silt and clay than is required for *loamy sands* are classed as *sands*. The humus content, as well as the clay and silt content, of *sands* is frequently very low; this characteristic distinguishes them in part from *loamy sands*.

A comparatively small percentage of clay imparts to a soil the characteristics of clay to a sufficient extent to warrant naming the soil a *clay loam*. For example, a soil sample containing equal proportions of sand, silt, and clay (point *B*, Fig. 16) is a *clay loam*. To persons accustomed to working in sandy loams, a clay loam seems very plastic when wet and hard when dry. Many farmers and others unfamiliar with the correct textural grouping of soils call clay loams "clay soils." As with sandy loams there are *heavy* and *light clay loams* in which the proportion of clay is near the upper and lower limits, respectively.

A soil containing enough clay to classify it in the clay loam group and, in addition, containing nearly the maximum quantity of silt is often very dense as well as plastic, and moisture moves through it slowly. However, such a soil, texturally classified as a *silty clay loam*, may be highly productive when skillfully tilled.

*Clay soils*, like silty clay loams, may be productive when in good physical condition, but they require special management methods to prevent the puddling or breaking down of the granules.

*Silt loams* lack the cohesive properties of clay and the grittiness of sand. Silt must be present in large quantities to impart its velvety or flour-like properties to a soil. Although silt loams are water-retentive, they lack the cohesiveness of clay loams and are accordingly easier to keep in good tilth.

**A Generalized Grouping of Soils.** The various soil classes are separated from one another by definite lines of division in Fig. 16. However, their properties do not change abruptly at these boundary lines,

but one class grades into the adjoining classes of lighter or heavier texture. For example, a loam gradually merges into a silt loam or clay loam as the percentages of silt and clay increase, and, on the other hand, a sandy loam is produced as the percentage of sand separates replace silt and clay in the loam. It is difficult, therefore, for those not well acquainted with the specific properties which characterize the different soil classes to distinguish accurately between them, and hence a more generalized textural classification of soils is sometimes used. For instance, it is customary, when speaking in general terms, to refer to loams and silt loams as medium-textured soils, to clay loams, silty clay loams, and clays as heavy soils, and to sandy loams, loamy sands, and sands as light soils. An area made up of heavy soils is frequently called *clay land*.

**Class Name Descriptive of Soil Properties.** It should be remembered that a soil class name is meant to convey not only an idea of the percentages of the various separates present but also *information concerning the soil properties associated with such a mixture*. Thus the ideas concerning tillage practices, moisture-retaining power, aeration, crop adaptation, and so forth, brought to mind by the term sandy loam should be quite different from those conveyed by the class name clay loam. However, these soil properties are also influenced by the mineralogical composition of the clay fraction and the structure of the soil.

### SOIL STRUCTURE

The term *texture* is used in reference to the size of soil particles, but, when the arrangement of the particles is being considered, the term *structure* is used. The structure of the different horizons of a soil profile is an essential characteristic of the soil just as are color, texture, or chemical composition. Structure influences moisture relations, availability of plant nutrients, action of microorganisms, and plant growth. Types of soil structure and the factors which influence the formation of them are suggested by the following questions.

#### Questions

1. What is meant by single-grained structure?
2. What forms of compound soil structure are recognized?
3. By what process are secondary soil particles developed?
4. How do moisture and temperature conditions influence extent of soil granulation?
5. In what ways does plant growth influence granulation of soils?

6. Do microorganisms have a part in aggregate formation?
7. How do wetting and drying affect the compound structure of soils?
8. Do freezing and thawing affect soil aggregation?

**Single-Grained Structure.** If each particle in a soil functions as an individual, that is, if it is not attached to other particles, the structure is called single-grained and the individual particles are known as primary particles. Such a structural condition may exist in very sandy soils or in the eluviated horizon of podzolic soils from which the clay and organic matter have been leached. Some of the particles in many soils exist in the single-grained condition, but to find all the particles so functioning is unusual. If soils containing considerable clay are tilled or are tramped by grazing animals when too wet, a portion of the clay particles assume the single-grained condition.

**Compound Structure.** When particles adhere in clusters and the clusters then function as particles, a compound structure is developed. These clusters or compound grains may be called secondary particles. Some of the very small particles combine into clusters too small to be seen by the naked eye and so may be said to develop a microstructure. The small units together with primary particles unite into larger and visible units of various shapes and sizes to develop a macrostructure. In the last few years much study has been devoted to soil structure and particularly to the micro units. The terms granules and aggregates are used to designate compound or secondary structural units without regard to size or shape.

Various systems have been suggested for classifying the macrostructural units of soil, other than primary particles, but none has been universally accepted. The following system is as useful as any and sufficiently detailed for most purposes.

- I. Particles or aggregates with all axes of approximately the same length, somewhat spherical or cubical in shape but exposing many rather than few faces, some of which are rounded.
  - a. Nut-like structure—units of considerable size, greater than 6 mm. in diameter.
  - b. Granular structure—units of smaller size, from 6 to 3 mm. in diameter.
  - c. Crumb structure—units are small, porous aggregates which are quite easily crushed.
- II. Particles or aggregates in which the vertical axis is appreciably longer than the others.
  - a. Prismatic structure—units with faces moderately to well defined.
  - b. Columnar structure—units are prismatic but have rounded tops.

- III. Particles or aggregates in which the horizontal axes are appreciably longer than the vertical.
  - a. Platy structure—units decidedly flattened but yet of considerable thickness.
  - b. Laminar structure—units quite thin.
- IV. Particles or aggregates with no uniformity in length of axes, faces well-defined but irregular in number and form.
  - a. Blocky structure—units of considerable size, greater than 1 cm.
  - b. Fragmental structure—units of smaller size, less than 1 cm.

**Formation of Secondary Particles.** An understanding of the causes for the development of compound structures in soils is of practical importance because structure has a great influence on plant growth and is also an important factor in soil classification.

Evidently there must be some agent which draws the particles together in order to produce aggregates and also some means by which they are bound rather firmly so that the structural forms may persist. Evidence points to the colloidal fraction as the active constituent in compound structural units, since without its presence single-grained structure prevails. It has been shown that there is a high degree of correlation between the amounts of clay (5 microns) in soils and the percentage of granules with a diameter greater than 0.05 mm. A similar correlation was found between percentages of aggregates of the same dimensions and the percentage of organic matter. Furthermore, a high correlation between organic content and development of aggregates larger than 0.1 mm. in diameter indicates that organic matter leads to formation of large granules. Three groups of colloidal matter are thought to serve as cementing material in aggregate formation: (1) clay minerals, (2) colloidal oxides of Fe and Al, and (3) colloidal organic matter.

There are several theories concerning the process by which aggregation is brought about, but the following is probably as widely accepted as any. Because colloidal particles are charged bodies, water molecules attach themselves readily and firmly to them. As water molecules and colloidal nuclei carry both positive and negative charges, it is conceivable that oriented water molecules may connect colloidal particles. The linkage may include adsorbed cations because aggregation seems to be affected by the cations adsorbed.

Such water molecules are held very tightly, and as water evaporates from the soil the number of water molecules attached to two particles increases, and at the same time the length of each linkage becomes shorter and stronger and so pulls the colloidal particles close together.

Larger soil particles to which the colloids are attached may be drawn into the cluster, and some may be surrounded as the cluster or floccule develops. As more water is lost and the colloidal material becomes further dehydrated, it sticks or cements the particle group into a unit which is permanent until rehydration of the colloid reduces its binding power. Thus it is seen that water, acting in conjunction with the colloids, constitutes the force which produces granulation of soil and that colloidal material is the final binding agent.

Many soluble salts also exert a strong flocculating effect on colloids, and hence their presence may be a factor in inducing granulation in soils. Applications of lime to very fine-textured soils in poor physical condition may noticeably increase granulation.

These processes of soil aggregation are greatly facilitated by other factors and processes which help to create lines of weakness in a soil mass and so permit groups of soil particles to be drawn around a new nucleus and finally cemented into a structural unit. In the following paragraphs some of the agencies which aid in soil granulation are discussed.

**Influence of Climate on Aggregation.** A relationship between the climatic conditions which have contributed to the formation of the great soil groups is discussed in Chapter 4. Moisture conditions which have produced the dark soils of the grasslands of the eastern Great Plains and the upper Mississippi Valley have led to a higher degree of aggregation than exists in the drier soils to the westward or in the forest soils to the east. The explanation lies in the fact that moisture supply was adequate for the development of considerable clay through mineral decomposition but was not sufficient to eluviate the clay as in the timbered soils. In the drier soils too small an amount of clay was developed to promote extensive aggregation. The large amount of organic matter in the prairie soils also promoted the development of a compound structure. The high organic content and lack of eluviation of the humid prairies led to a high degree of aggregation in these soils.

Low temperatures also promote the development of compound soil structures, as is evidenced by the much higher degree of aggregation in the northern than in the southern Great Plains grasslands. The explanation of this situation probably lies in the greater content of organic matter accumulated under lower temperatures when moisture conditions are relatively constant. Furthermore, the northern grasslands are submitted to frequent freezing and thawing. A reversal of the effect of temperature on degree of aggregation is found in the soils of the humid region in that the soils of southeastern United States (lateritic soils)

show a high aggregation of the silt and clay (about 75 per cent). This is due to the binding power of the dehydrated oxides of iron and aluminum. The forest soils of the northern humid region show a much lower development of compound structure because of eluviation of clay and organic colloids, as previously mentioned.

**Granulation Induced by Plant Roots.** It has long been observed that vegetative cover influences the development of compound structure in soils. Grass appears to be the most efficient type of vegetation in inducing aggregation, probably because of the completeness with which its root system fills the soil. Several means by which plant roots cause aggregation have been suggested. They may be summarized as follows: (1) Roots and root hairs penetrating the soil produce lines of weakness along which the clod or soil mass may break into granules. (2) The pressure exerted by developing roots may induce aggregation. (3) Root secretions may flocculate colloids, stabilize, or cement aggregates. (4) Use of moisture by roots may cause dehydration of colloids, thus resulting in shrinkage and finally in cementation. (5) Organic colloids produced from decay of roots may be an aggregating agent.

**Microbial Activity and Aggregate Formation.** Many of the observed effects of soil organic matter on aggregation of soil particles can be attributed to microbial activity. The growth of microorganisms leads to a binding together of soil particles and, consequently, an increase in soil aggregation. For example, it has been found that fungi not only cause a mechanical binding of particles due to extensive mycelial growth but also bring about aggregation as a result of synthesizing certain organic compounds. Thus, in addition to the direct mechanical effects of mycelia, the suggestion has been made that the soil microbes produce gums, waxes, and possibly other water-insoluble substances which coat the soil particles and increase aggregation.

**Wetting and Drying Influence Structure.** The granulating effect of the wetting and drying of soil is familiar to all who work with or till the soil. The disintegration of clods into granules under the influence of alternate wetting and drying is a common observation. The explanation of the process probably lies in the alternate swelling and shrinking of the colloids. Uneven expansion and contraction, which must occur because a portion of soil is never wet or dried uniformly and simultaneously, result in numerous cracks which separate the soil into smaller pieces. They may be drawn apart into units as shrinkage of the colloid occurs.

A contributing factor in disrupting a fragment or clod is the com-

pression of the air trapped in the interparticle spaces as the exterior of the clod is wet. As the moisture moves inward, the pressure becomes greater until finally the compressed air explodes the clod into fragments.

**Aggregation Induced by Freezing and Thawing.** Aside from the desire to get work out of the way in order to lighten the spring work schedule, farmers explain their interest in fall plowing on the basis of the improved tilth which it encourages in heavy soils. This increased granulation is due in part to repeated wetting and drying and in part to alternate freezing and thawing which take place much more readily in plowed land. Plowing also relieves the lateral pressures which oppose the granulating effect of the processes described.

As soil water freezes in a small crevice of the soil, it draws moisture from the surrounding soil as the ice crystals grow. Thus, under slow freezing the ice crystals tend to break the clod and at the same time exert a pressure on the surrounding soil particles which presses them together and stimulates the development of new clusters or granules. Furthermore, as moisture is drawn from the adjoining soil, the colloidal material is partially dehydrated and shrinks and tends to cement the particles into an aggregate. The withdrawal of moisture sometimes also leads to the concentration of soluble salts which flocculate the colloids, and thus the foundation is laid for soil granulation. Repetitions of freezing and thawing from night to day accentuate these processes until they culminate in a thoroughly granulated soil. The role of water in this process is shown by the fact that freezing and thawing of relatively dry soils do not induce granulation. Also, it has been demonstrated that very quick freezing splits up soil aggregates instead of inducing granulation. This result comes about through the formation of many small crystals because there is not time for sufficient water to move to zones of crystal formation to make large crystals.

#### TILLAGE OPERATIONS AND SOIL PROPERTIES

The texture of a soil can be altered by man only through the addition of soil separates, as is sometimes done by applying sand to clay soil or clay soil to sand. In irrigation sufficient quantities of silt are sometimes added in the water to alter appreciably the texture of the soil. Man spends much time, however, in improving the structure of the soil. The plow is the most effective farm implement for altering soil structure. It has been estimated that about one-third of all draft power used on farms is expended in plowing. Virtually all tillage practices, except those directed toward the killing of weeds, are carried out for the purpose of developing a desired soil structure.

If tillage operations are to improve soil structure, they must be carried on when the soil holds a suitable moisture content. When the soil is too moist, the cementing material of the granules is so soft that the granules are destroyed. The particles readily slide over one another, smaller ones slipping in between larger ones to create a compact mass dominated by a single-grained structure. The soil is then said to be puddled. On the other hand, if the soil is too dry when tilled, it breaks up into large lumps. To restore a heavy soil, which has been worked when too wet or too dry, to a desirable structural condition frequently requires much subsequent tillage combined with wetting and drying or freezing and thawing. A year or several years may be necessary to undo the harmful effects. Some tillage implements and tillage operations are discussed under the headings suggested by a series of questions.

### Questions

1. How was the plow developed?
2. How does the plow induce soil granulation?
3. What types of plows are there?
4. What are the comparative merits of fall and spring plowing?
5. How deep should one plow?
6. For what purposes are harrows used?
7. What are the functions of rollers?
8. Why are row crops intertilled?
9. Is excessive tillage of the soil undesirable?

**History of the Plow.** Some type of plow for the purpose of loosening or turning over the soil was probably among the first tillage implements used. Paintings on the walls of ancient Egyptian tombs, dating back 5,000 years, depict oxen yoked together by the horns, drawing a plow made from a forked tree. The Greeks improved the Egyptian plow by adding a metal point. There was little additional improvement in the plow until American Colonial days, when a wooden-framed plow, having the moldboard covered with iron strips and carrying a curved iron point or share, was constructed.

Thomas Jefferson, who made the first scientific study of the plow, considered that it consisted of two wedges, one to cut the furrow slice free at the bottom and lift it and the other to cut the soil at the side and turn it. He calculated a correct mathematical combination of the two wedges. The cast-iron plow was made about 1800.

**Granulating Action of the Plow.** The design of the moldboard plow has been carefully worked out and the slopes and curvatures of its

various parts calculated in order to produce the desired effect on the soil without requiring an excessive amount of power. As the furrow slice is cut free and passes over the sloping share, it is lifted and pushed sideways so that it is submitted to a vertical and horizontal strain which tends to break it into layers perpendicular to the line of movement of the plow. At the same time a strain parallel to the passage of the plow is developed, owing to the friction between the soil and plow bottom. This stress results in a shearing of the soil into layers which are perpendicular to those previously mentioned. The result is that the furrow slice is fractured in three directions. As it passes over the curved moldboard, these fractures are amplified and the soil is thrown over in a completely shattered state so that the forces of granulation may draw the particles together and cement them into definite units.

**Types of Plows.** There are several types of moldboard plows, the most common of which are the (1) stubble, (2) general-purpose, and (3) sod, which are different in length and in the curvature of the moldboard. The stubble plow has a short, highly curved moldboard and hence accomplishes the maximum of soil pulverization. It is used primarily in light-textured soils and in those which are infrequently planted to grasses which bind the soil together by a dense growth of roots. The sod plow is used for breaking heavy sods in which the soil is permeated with roots and breaks up with great difficulty, the furrow slice tending to hang together as a continuous unit. This plow has a long, gently curved moldboard. The use of a sharply curved

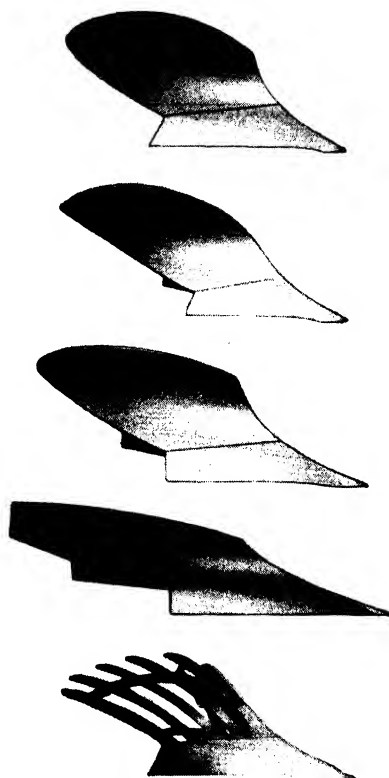


FIG. 17. Different types of plow bottoms showing variations in length and curvature of the moldboards. From top to bottom: stubble, blackland, general-purpose, prairie breaker, and slat moldboard. [Courtesy of Deere and Company, Moline, Illinois.]

moldboard in such soil would require a great deal of power and would not accomplish desirable results. The general-purpose plow is intermediate, in length and curvature of moldboard, between the two described. It is the most commonly used plow in general farming on the loam and heavier soil classes.

The lister is essentially a double plow with right- and left-hand

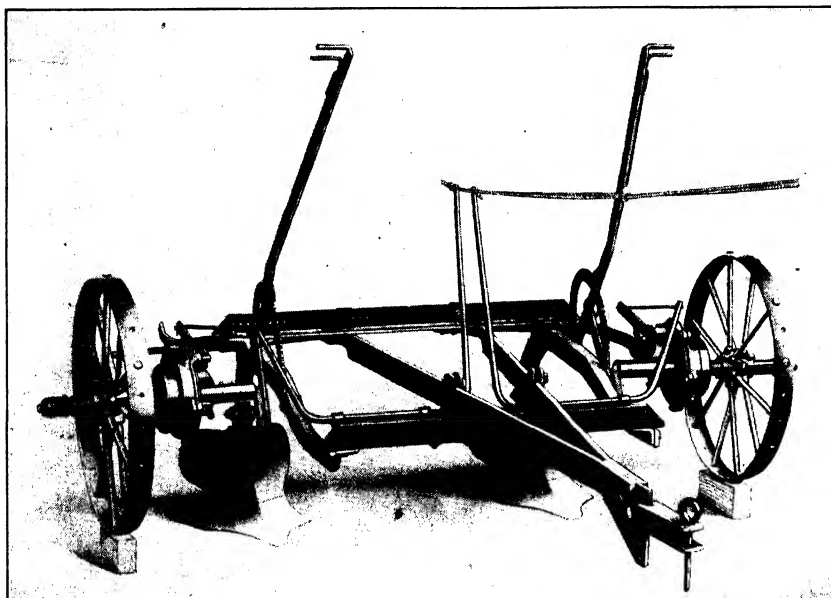


FIG. 18. The lister has a double moldboard so that it turns a furrow slice in both directions, thus forming a broad trench. The seed of grain crops planted in rows is often planted in the bottom of the trench. [Courtesy of J. I. Case Company, Racine, Wisconsin.]

bottoms mounted back to back. Hence it throws a furrow slice in each direction. The resulting ridges frequently are split, thus turning over the entire soil. The lister is used for ridging ground in the growing of special crops. On the Great Plains the lister is used for planting corn and various sorghums, the planting device being a part of the plow and dropping and covering the seed in the bottom of the trench. Subsequent cultivations throw the soil from the ridges around the growing plants. Lister furrows, plowed at right angles to the prevailing winds, materially decrease soil blowing.

Various types of moldboard plows with either two bottoms or a reversible moldboard are made for use in hilly country or on experi-

mental plots where it is desired to throw all the furrows in one direction.

The disk type of plow is convenient for use in hard or dry soils that are not easily penetrated by the moldboard plow and in loose, unconsolidated sandy soils. Sticky soils in which a moldboard will not scour

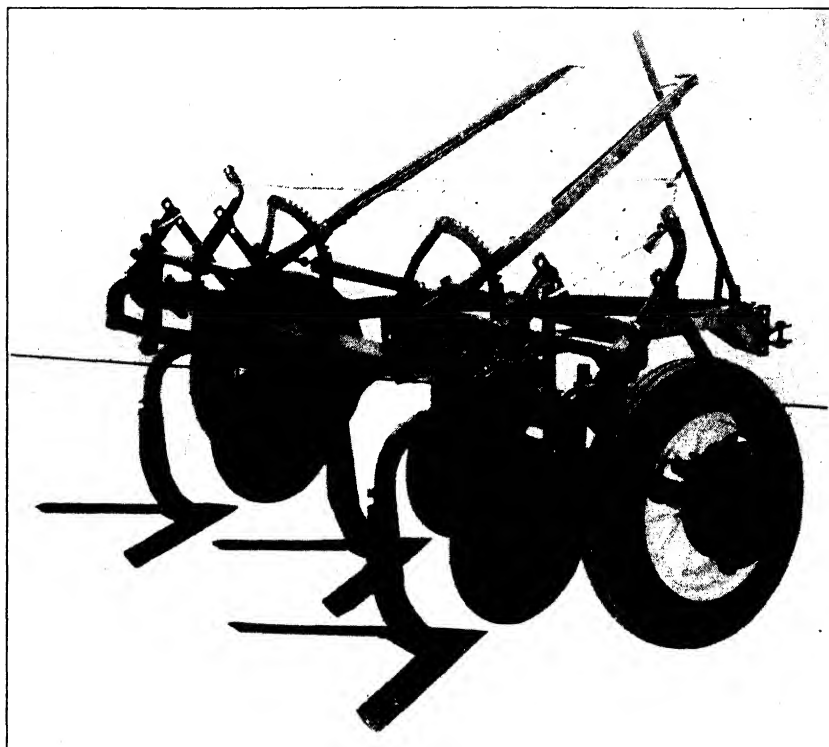


FIG. 19. Sweeps are used to cut the roots of growing plants and to lift and loosen the soil without turning it over or working into the soil crop residues which are on the surface. [Courtesy of Deere and Company, Moline, Illinois.]

and soils containing many larger roots and shrubs are more easily plowed with a disk than with a conventional type of plow. The straightness with which the disks are set and the adjustment of scrapers at the top of the disks determine the extent to which the furrow slice is turned over. This type of plow is used extensively in some parts of the country, especially in the Great Plains. Its simplicity and ability to operate under a wide variety of conditions contribute to its popularity. On the other hand, it tends to "fine" the soil more than is always desirable. The

implement is known by many names some of which are "one-way," "disk tiller," "harrow plow," "wheatland plow."

In areas of limited rainfall an implement which has broad blades set at an angle and fastened together to form a V is used. These blades pass through the soil at a depth of several inches, loosening it and cut-



FIG. 20. An implement with chisel-like points mounted on strong curved bars loosens the soil and leaves crop residues largely on the surface as a protection against wind and water erosion. [Courtesy of Graham-Hoeme Plow Company, Amarillo, Texas.]

ting off the roots of weeds, grass, or other plants. The soil is not turned over, however, and neither are plant residues on the surface incorporated in the soil to an appreciable extent. Implements of this type are usually designated "sweeps."

Another implement that is meeting with considerable favor, especially in the South, consists of a series of chisel-type blades on strong curved bars. These are staggered to give complete loosening of the surface soil without inverting the plow slice. The chisels can be set to penetrate from 1 or 2 to about 8 inches. The implement is of particular value in

a mulch type of culture where crop residues are left on the soil surface. Heavy soils that get very hard may be loosened with this machine but usually require disking also to prepare a seedbed. Sandy soils may require no additional tillage before planting.

**Fall vs. Spring Plowing.** Fall plowing in some places is something of a habit and a tradition with farmers. In some agricultural sections of the country it was a custom to rate a farmer by the amount of fall plowing he was able to finish, and to some extent the same sentiment prevails today. When horses furnished the power on farms, plowing was a slow process and fall plowing was necessary if the spring work was to be done in time. The tractor has changed this situation, and plowing which formerly required several weeks may now be done in a few days. With the passing of the horse has gone much of the necessity for fall plowing; yet it is practiced to a far greater extent than is good for the land. Fall plowing opens the way for loss by leaching of soluble plant nutrients, especially nitrates and lime. Fall plowing prevents the use of winter cover crops, which decrease erosion and the leaching of nutrients and add organic matter to the soil. And, of more importance, fall plowing of all but very level land encourages loss of soil by erosion, which often results in greater soil depletion than does crop production.

The beneficial effects on soil structure of plowing heavy land in the fall have been previously discussed. They cannot be ignored on heavy, sticky soils, particularly those that are level. Furthermore, spring grain usually may be seeded earlier on heavy soils which have been fall-plowed. But, for crops planted in the late spring or early summer, spring plowing is just as satisfactory on the majority of soils. Many farmers experience unsatisfactory results from spring plowing because of insufficient preparation of the soil after plowing.

**Depth of Plowing.** Many comparisons have been made of different depths of plowing. In most cases little advantage has been found from plowing to a greater depth than 6 to 7 inches. Some benefit from deeper plowing of very compact soils may result occasionally, particularly in the growing of root crops. Shallow plowing has also been found desirable on some occasions when planting must follow so shortly after plowing that there is not time to prepare properly the deeper-plowed soil.

**Harrows.** With the exception of the disk, various types of harrows are designed primarily to stir the surface soil to a depth of 2 or 3 inches, thus killing weeds, breaking clods, leveling the surface, and drying out the surface of the plow layer by exposing it to the air. The harrow com-

pacts the soil very little but may aid appreciably in inducing granulation. Numerous types of harrows are described in any good book about tillage implements.

The disk harrow has diverse uses. With the gangs turned at a sharp angle the soil is moved considerably; thus it is loosened and granulation is facilitated. Manure or other organic matter can be incorporated into the soil by means of the disk. Use of the disk harrow before plowing cuts up crop remains on the surface and mixes them with the soil so that more complete mixing with the plow layer may be accomplished by plowing. In hard or compact soil the surface is also loosened, permitting the furrow slices to settle better and making quicker and more complete contact between plowed layers and the soil beneath. Indeed, on many soils, unless there is considerable organic matter to work into the soil, a satisfactory seedbed for most small grains can be prepared by disking in place of plowing.

**Rollers.** Rollers are used to press down the furrow slices after plowing, to crush clods, to compact the soil around recently sown seed, to firm the soil around the roots of plants which have been loosened by heaving, and to settle and compact soil in seedbed preparation. The smooth roller has given place largely to those of the corrugated type because the corrugated roller penetrates the soil deeper, thus giving more subsurface packing. Corrugated rollers exert a side as well as a downward pressure and leave the soil in ridges rather than smooth, a desirable feature. In some types the rear gang may be raised, thus increasing the weight on the front gang.

**Intertillage of Crops.** Numerous types of cultivators are available for use on different kinds of soils with different crops and under various climatic conditions. There are four primary purposes for which the intertillage of crops may be practiced: (1) to destroy weeds, (2) to conserve moisture, (3) to increase the absorption of moisture on sloping land, especially on soils which tend to bake or crust at the surface, and (4) to throw soil to or away from the plants as deemed advisable.

It is generally customary to make the first cultivation deeper than those that follow in order to loosen soil which later will be filled with roots and hence should not be disturbed. Later cultivations, and sometimes the first one, are made with a blade or with some type of broad flat shovel (such as the duckfoot) which stirs the soil to a shallow depth only, thus preventing the cutting of roots.

The most important objective in intertillage is the elimination of weeds that compete with crop plants for moisture and plant nutrients.

After weeds are eliminated so that they do not use moisture, further cultivation generally conserves little more moisture on many soils. Considerable moisture may be prevented from evaporating, however, from soils which are very compact or heavy-textured or which tend to crust and crack badly. This point is discussed more fully in Chapter 8.

Many soils tend to crust on drying, allowing rain to run off instead of entering the soil. The granules in the surface of some soils are readily destroyed by rain, especially a beating rain, and the soil particles then pack together tightly, making a coating over the soil and permitting much of the water falling during the rest of the storm to run off. To overcome these tendencies and to roughen the surface and so impede the runoff of water, it is desirable to keep soils with an appreciable slope loosened and if possible somewhat ridged at right angles to the slope.

For crops planted or drilled in a continuous row, thus permitting cultivation in one direction only, it is desirable to throw a certain amount of soil toward the row to smother weeds between the plants. Soil is also thrown toward crops planted in lister furrows in order to conserve moisture and kill weeds. Some root and tuber crops need to be covered by soil as they develop in order to prevent sunburn and sometimes frost damage.

Intertillage or cultivation of the soil during the growth of a crop is considered advisable in some soils in order to hasten the decay of organic matter by microorganisms and thus produce nitrates and liberate other nutrients for use by the crop. The necessity for stirring the soil to increase the action of microorganisms varies with different soil and climatic conditions.

**Adequate but Not Excessive Tillage.** Sufficient tillage should be given a soil to prepare a suitable seedbed and to destroy as many weeds as possible before the crop is planted. It is commonly considered poor economy to slight the preparation of the seedbed. Subsequent tillage cannot compensate for the handicap imposed on crops by planting them in a soil insufficiently prepared. For most crops the soil should be thoroughly settled and firm to the depth of plowing. There should be only enough loose soil to insure planting at a proper depth with good coverage of the seed. In regions of limited rainfall, seedbed preparation should cover a suitable period and include such tillage as is needed to provide for the storage of soil moisture. On the other hand, there is a growing sentiment favoring plowing shortly before planting with little subsequent tillage in order to bury germinating weed seed, reduce labor,

and not damage soil structure by excessive tillage and use of heavy equipment.

After planting and before the plants have emerged or attained too great a height, many weeds may be destroyed with a harrow at a minimum of cost. Later tillage should be confined to what is necessary to attain the objectives set forth in the preceding section. More tillage not only adds needlessly to the cost of crop production but also is detrimental to the soil because it results in excessive decay of organic matter. Although some farmers may not fit good seedbeds and cultivate their crops adequately, there are many who stir the soil to excess with tillage implements. Plowing when it is not necessary for seedbed preparation and excessive intertillage of crops are detrimental to soils.

#### POROSITY AND WEIGHT OF SOIL

That portion of a given volume of soil which is unfilled with solid matter is termed *pore space*. The proportion of a soil occupied by pore space depends on both the texture and structure of the soil and on the shape of the particles. A high content of organic matter also leads to much pore space. In soils containing little silt and clay the total amount of pore space is small even though the individual spaces are large. In soils containing a high proportion of silt and clay the relative volume occupied by pores is large, and if these soils are well granulated the total volume of pore space may be equal to or even greater than that occupied by the solid particles. This is because space exists not only between soil grains but also between granules.

The amount of pore space in a soil is expressed as a percentage of the total soil volume. The term *porosity* refers to the total pore space in a soil rather than to the size of the individual pores.

Pore spaces are important because they contain air and moisture, both of which play just as much a part in the crop-producing capacity of a soil as the mineral and organic matter. The amount of pore space in a given volume of soil greatly influences its weight. Several questions concerning the porosity and weight of soil and their interrelationships are listed.

#### Questions

1. What is meant by the specific gravity of soil?
2. How does volume weight differ from specific gravity?
3. How much does an acre of soil, to a depth of 6½ inches, weigh?
4. How is porosity determined?

**Specific Gravity of Soil.** In determining the specific gravity of soil, consideration is given to the solid particles only. Thus, the specific gravity of any soil is a constant and does not vary with the amount of space between the particles. Specific gravity is determined by dividing the weight of a given volume of dry soil particles by the weight of water that these particles will displace. Specific gravity is a weight ratio and does not vary a great deal for different soils unless there is considerable variation in content of organic matter or mineralogical composition. It averages about 2.65.

**Volume Weight of Soil.** In determining volume weight of soil, consideration is given to the pore space as well as to the solid particles. Volume weight of a given soil is, accordingly, a variable because the volume of pore space varies. It is calculated by dividing the dry weight of a given volume of soil in its natural structural condition by the weight of an equal volume of water. In determining the volume weight of field soils, great care is exercised not to destroy their structure. Samples are taken by driving into the soil a steel tube with a sharp cutting edge. The inside diameter of the tube is increased slightly immediately above the cutting edge so that the core of soil may be readily removed. The outside of the tube just above the cutting edge is also enlarged so that the soil is pressed outward as the tube is driven in, thus enlarging the hole and making withdrawal of the tube easy. As the dimensions of the tube are known, the volume of the soil core removed is known, and so by drying and weighing it the necessary data are obtained for calculating the volume weight. *Apparent specific gravity* is a term used synonymously with *volume weight*. The volume weight of heavy soils may range from 1.1 to 1.6 and of sandy soils from 1.3 to 1.7.

**Weight of Field Soils.** As volume weight varies so greatly with soil structure, so the weight of soils under field conditions must vary. A cubic foot of loam, sandy loam, or sandy soil in the dry condition weighs 80 to 110 pounds, whereas a cubic foot of silt loam, clay loam, or clay ranges from 70 to 100 pounds in weight. It is generally estimated that an acre of the finer-textured soils to the average depth of plowing (about 6½ inches) will weigh approximately 2,000,000 pounds when dry. This weight is used frequently in reporting the content of plant nutrients or of organic matter in terms of pounds per acre. Sandy and sandy loam soils have a somewhat greater weight. The weight of soils under field conditions, that is, with normal moisture content, is much greater. As a result, in plowing an acre a very appreciable weight is turned over.

**Determining Porosity of Soil.** The percentage of pore space in a soil may be calculated from the volume weight and specific gravity. As specific gravity is assumed to be about the same for most soils, a volume-weight determination is all that is required for the calculation of pore space. If it is borne in mind that both specific gravity and volume weight are based on the weight of unit volume of water (1 ml. = 1 gram), it is readily seen that (volume weight ÷ specific gravity) × 100 gives the percentage of a volume of soil that is filled with solid particles. This percentage, taken from the total volume (100 per cent), will give the percentage of pore space; hence the formula

$$100\% - \left( \frac{\text{volume weight}}{\text{specific gravity}} \times 100 \right) = \% \text{ pore space}$$

### SOIL COLOR

The color of soil is a factor of service to both farmer and soil scientist, provided that one understands the causes of the various colors and is able to interpret them in terms of soil properties. Organic matter content, drainage condition, and freedom of aeration are soil properties related to color which are of interest to farmers. The investigator uses color as an aid in soil classification and draws from the color of the different horizons information about conditions pertaining to and forces active during soil formation. The following division of the subject matter may be made.

#### Questions

1. What is color?
2. What are the standard conditions for observing soil color?
3. How can the qualities of light causing the sensation of color be measured?
4. Can the measurement of color be translated into terms which convey the proper impression to others?
5. What are the names of these different colors?
6. What are the colors of soil minerals?
7. What causes gray and brown colors in soils?
8. Why are some soils red or yellow?

**Color Produced by Reflected Light.** Color is the sensation produced when light from an object enters the human eye. Thus the color of a soil is the result of light reflected from the soil. Consequently, in order to describe it concisely it is necessary to define carefully (1) the

qualities of the light falling on the soil; (2) the soil conditions affecting the reflection of this light; and (3) some standard method for measuring the qualities of this reflected light. Then finally we need some system of nomenclature to translate these measurements into words that will convey our observations to other persons.

**Standard Conditions for Observing Soil Color.** The usual standard light source for the observation of soil colors is the light from a clear sky. This closely approaches white light; that is, all visible wavelengths of light are present in nearly constant proportions.

A freshly broken soil surface or a natural structure face is usually chosen for observations. The color of this sample should be recorded at two standard moisture contents: (1) at field moisture capacity, and (2) when the sample is air-dry. It is usually more practical in field work to use moist samples, because a sample can be moistened and then the color observed as soon as the visible moisture films have disappeared from the surface. However, it is frequently impractical to wait for a moist soil to dry out. Soil colors generally become whiter on drying, but they vary greatly in the magnitude of the color differences at different moisture contents. Consequently, color descriptions are most helpful when made under both of the suggested moisture conditions.

**Measurement of the Quality of Light.** The color of light can be accurately described by measuring its three principal properties, called hue, value, and chroma. Hue refers to the dominant wavelength or color of the light. Value, sometimes called brilliance, refers to the total quantity of light. It increases from dark to light colors. Chroma is the relative purity of the dominant wavelength of light. It increases with decreasing proportions of white light.

**Color Designations.** The Munsell notation of color is a systematic numerical and letter designation of each of the three variable properties of color. The relationships of the colors to one another can be shown by use of a solid, for example a cube in which hue, value, and chroma are plotted along the three edges. Each possible color represents a point in this cube and is completely defined by the three coordinates of that point, which is its Munsell notation. The three properties are always given in the order hue, value, and chroma. For example, in the Munsell notation 1Y 6/4, 1Y is the hue, 6 is the value, and 4 is the chroma.

The Munsell notation for a given soil sample can be quickly determined by comparison of the sample with a standard set of color chips. The chips are mounted in a notebook with all the colors of a given hue on one page. Each page then corresponds to a slice through the color cube parallel to its front. The pages are arranged in the order of in-

creasing or decreasing wavelength of the dominant color in order to facilitate the matching of the unknown soil color with the color of the standards.

**Color Names.** Because there are an infinite number of points in the color cube, it is impossible to have a common color name for each possible Munsell notation. However, names can be used for colors having a relatively narrow range in each of the three properties of color. Several systems of color names have been proposed, two of which are commonly used in the United States: (1) The ISCC-NBS system, proposed by the Inter-Society Color Committee in cooperation with the National Bureau of Standards, defines common color names that are intended to apply to the colors of all objects; (2) the system proposed by the Soil Survey Division of the U. S. Department of Agriculture is for describing soil colors. It would be helpful if a uniform system of color names could be adopted. The Soil Survey Division follows the practice of giving the soil color name and also the Munsell notation of that color in soil series descriptions. For example, the Munsell notation 1Y 6/4 is a light yellowish brown.

**Colors of Soil Minerals.** The minerals occurring in appreciable quantities in most soils are light in color. As a result, soils would be of a light gray color if composed of crushed minerals which had undergone little chemical change. Accordingly, for an explanation of the dark gray, brown, red, and yellow colors in soils we must look to chemical changes in the constituents (especially iron) of the minerals and to the addition of organic matter. There are a few instances, however, in which the proportion of colored minerals is sufficient to give the soil a decided color.

**Gray and Brown Soils.** The dark color of soils is generally due to the highly decayed organic matter they contain; in fact, with some practice the percentage of organic matter in a soil may be judged with reasonable accuracy from its color. Although we speak of black soils, there are none which are truly black. Organic matter imparts a gray, dark gray, or dark brown color to soils unless some other constituent such as iron oxide or accumulation of salts modifies the color.

If soils are poorly drained there is usually a greater accumulation of organic matter in the surface layers, thus giving a very dark color. The lower soil layers, which contain very little organic matter, on the other hand, are of a light gray color indicating the poorly drained condition. If drainage is intermediate, the gray of the subsoil is likely to be broken by flecks of yellow.

**Red and Yellow Soil Colors.** When drainage permits aeration, and moisture and temperature conditions are favorable for chemical activity, the iron in soil minerals is oxidized and hydrated into red and yellow compounds. Highly hydrated iron oxides are yellow, but as hydration diminishes reds replace the yellows. Accordingly, we find shades of red in soils extending from the southern deserts across the semi-arid and subhumid belts to the humid states of the Southeast. The red and yellow colors of the subsoils in the Southeastern States immediately catch the eye of the traveler. The low organic matter content of many soils in this area leaves undulled the brilliant colors of the iron oxides. With an appreciable humus content the red colors are converted into mahogany colors.

### SOIL TEMPERATURE

As temperatures decrease, the life processes of both plants and animals are slowed down until finally they cease altogether. Growth processes of most agriculturally important plants are very sluggish at temperatures about 40°F. and increase until temperatures ranging from 70° to 90°F. are reached. The chemical processes and activities of microorganisms which convert plant nutrients into available forms are also materially influenced by temperature. The farmer is much interested therefore in soil as well as in air temperatures, and a brief discussion of the topic follows.

#### Questions

1. What sources of heat warm the soil?
2. What factors determine the heat capacity of soil?
3. By what practices can man control soil temperature?
4. How does situation influence soil temperature?

**Sources of Soil Heat.** The movement of heat from the warm interior of the earth is counteracted by the loss of heat from the surface by radiation. Thus at a given depth, about 3 or 4 feet at the equator and 50 feet in the latitude of New York, the earth remains at an approximately uniform temperature, which is the same as the average annual air temperature, and the growth of plants is unaffected through the warming of the soil around their roots by the interior heat of the earth.

Likewise the soil is not warmed by the decay of organic matter within it, for the heat so generated is liberated so slowly that the effect on soil temperature is not measurable.

Man must depend, therefore, on the sun to warm the soil sufficiently to enable crops to grow. Even the radiant energy of the sun would not

perform this task satisfactorily if it were not for the tempering influence of the atmospheric envelope around the earth. Without the modifying influence of the atmosphere, the sun's heat by day would be too intense and at night radiation would proceed so rapidly that life would be destroyed through freezing.

**Heat Capacity of Soil.** Mineral particles require a comparatively small amount of heat to raise their temperature. The quantity of heat required to raise the temperature of a gram of soil particles  $1^{\circ}\text{C}$ . is only about one-fifth as much as is required to warm a gram of water the same amount. In other words, the specific heat of dry soil particles is 0.2, and it is evident from this fact that moisture content is the controlling factor in determining soil temperature. Soil high in water content will warm up slowly in the spring and will cool down slowly in the fall. Drainage therefore exerts a major influence on soil temperature.

**Control of Soil Temperature.** As has just been pointed out, the removal of excess water from a soil will facilitate changes in soil temperature. By providing drainage, man may exert some influence on the temperature relations of soils which are so situated that they hold excessive quantities of water. By use of mulches and various shading devices, the soil may be protected from the sun's rays and evaporation of moisture may be reduced, thus modifying changes in soil temperature. Aside from these practices there is little that man can do to affect the temperature of the soil he tills except to keep it in good tilth and increase the humus content. Soils high in organic content absorb more heat because of their dark color and hence tend to be warmer. But this effect is offset to some extent by the increased water-holding capacity. When in good structural condition, soils also warm up more readily than when they are in poor tilth.

**Location and Temperature.** In the northern hemisphere soils which are located on southern and southeastern slopes warm up more rapidly in the morning than if they were located on the level or on the northern slopes. The reason is that they are more nearly perpendicular to the sun's rays, and hence a maximum amount of radiant energy strikes a given area. Soils with a southern or southeastern exposure often are selected for the growing of early vegetables and fruits.

# 4

## Classification of Soils

The properties and the management of soils may be most effectively studied and remembered if the soils are classified according to a definite system. Several schemes for classifying soils have been suggested at various times, but one based on the properties of the soils themselves—rather than on kind of material from which they were derived, or on agency by which the material was accumulated, or on other factors having to do with soil development—appears most logical and advantageous. A system of this kind, called a *natural* classification, has been developed in the United States.

A *genetic* classification was suggested about 1880 by the Russian scientist Dokuchaev, and it has been further developed by European and American workers. The system is based on the theory that soils have a definite morphology (form and structure) which is related to a particular association of climate, vegetative cover, relief, age, and parent material. The classification of soils and the utilization of the soil groups by man are discussed under six general headings.

### Objectives

- A. A systematic classification of soils.
- B. Soil mapping and the soil survey.
- C. Relation between soil groups and climatic conditions.
- D. Age, relief, and parent material in relation to soil groups.
- E. Relation between soil groups and vegetative cover.
- F. Soil groups in relation to population density and the production of food and fiber.

### A SYSTEMATIC CLASSIFICATION OF SOILS

The modern system of classification is based on the recognition of soils as natural entities possessing definite unique characteristics. This scientific classification is often confused with better-known special systems that have evolved or have been devised from time to time to meet

specific requirements. Some examples of these special groupings are found in the classification of soils according to their lime requirements, according to their abilities to produce special crops, or according to their moisture-retaining characteristics. As new facts have been collected, the pedologic system gradually has grown. The growth of soil science necessarily has paralleled the accumulation of knowledge concerning soil relationships. Below are important questions to consider in a discussion of this subject.

### Questions

1. How may soil be defined pedologically?
2. Into what two broad groups are soils divided?
3. How are organic soils classified?
4. According to which characteristics are soils divided into series?
5. What information about a soil is conveyed by the type name?
6. Is there a similarity in the systems of classifying soils and plants?
7. What are the differences between the three soil orders?
8. What divisions are made in suborders of soils?

**A Pedologic Definition of Soil.** The noun *soil* is derived, by way of Old French, from the Latin *solum*, which commonly had the same meaning as our modern word. In general, soil refers to the loose surface of the earth as distinguished from solid rock, although many people, when they think of the word soil, have in mind that material which nourishes and supports growing plants. This meaning is even more general, since it includes not only soil, in the common sense, but also such things as rocks, water, snow, and even air—all of which are capable of supporting plant life. The farmer, of course, has a more practical conception of the meaning of soil; he says that it is the medium in which crops can grow. The civil engineer, on the other hand, looks upon soil as that material which supports foundations, roads, or sewers. In short, the word soil has many meanings. In talking or writing about soils, one must understand that the word can be used in these various ways.

By soil the soil scientist generally means that part of the earth's crust which has been changed as a result of soil-forming processes, that is, the *solum*. Pedology has adopted the old Latin root in order to make clear this definition of the word soil. A little more than just *solum* is intended, however, because the pedologist deals with areas, not merely single vertical profiles in a landscape. In discussions of pedologic topics like development and classification, soil always means either the *solum* or the profile of a soil type. In discussions of other subjects such as

fertility and productivity, soil is often used in the looser sense of something that supports plant life; in a vague manner, this meaning implies a definite profile, but it also implies to many people the concept of productive land. In reading this book, then, it is well to remember the different meanings that the word soil may express. Some students are often confused because they make no attempt to interpret the term in its appropriate sense.

**A Broad Grouping of Soils.** The first step in classifying soils is to divide them into two broad groups, namely, (1) organic soils and (2) mineral soils. Organic soils are defined as those which contain sufficient organic matter that the soil properties are dominated by the characteristics of the organic rather than by those of the mineral material. Rather empirically, the percentage of organic matter necessary to place a soil in the organic group is set at 20. In some states all organic soils are classed as *peat*. In other states the term *muck* is used to designate a soil in which the organic parent material is quite highly decomposed. And soils in which the parent material is yet fibrous and little decomposed are termed *peat*. The muck and peat soils belong to the Bog and Half-bog great soil groups in Chart 2 (p. 92).

**No Generally Accepted Classification of Organic Soils.** None of the proposed systems for classifying organic soils meets all requirements. Dachnowski<sup>1</sup> has proposed a simple grouping which includes (1) sedimentary or pulpy peat (derived from sediments and aquatic plants), (2) fibrous peat (high-moor and low-moor), and (3) woody peat (swamps and forest). Waksman<sup>2</sup> used a similar but somewhat enlarged grouping for the classification of New Jersey peats. The four classes include: (1) sphagnum or moss peat, (2) sedge and reed peat, (3) forest peat, and (4) alluvial or sedimentary peats. The last group is represented by salt-marsh peats which contain coarse fibrous material and large amounts of mineral sediments. It should also be noted that there is little sphagnum peat in the United States. Most of the moss peat is derived from *hypnum* and similar mosses.

Neither of the systems mentioned meets the need from the standpoint of utilization. The degree of decomposition of the organic material is important from the standpoint of crop production, as is also the degree of acidity. The kind of plants from which the deposit was derived may or may not be a major consideration but usually is of less significance than stage of decomposition and reaction. For the classification of

<sup>1</sup> *Soil Science*, Vol. 17, p. 107, 1924.

<sup>2</sup> *N. J. Dept. of Conservation and Development Bull.* 55.

organic soils in Michigan, Veatch<sup>3</sup> developed a system based chiefly on color, structure, stage of decomposition, content of mineral matter, and reaction. The general plan of the system is as follows:

- I. Older soils in advanced stages of decomposition.
  - A. Soils having little or no mixed mineral material.
    1. Dark brown or black soils high in lime.
    2. Brown soils, acid or less high in lime.
  - B. Soils containing a large proportion of mineral material.
    1. Soils in which the alluvium is admixed or in alternating layers.
    2. Clayey or silty mucks.
- II. Younger soils in which the organic matter is only slightly decomposed.
  - A. Soils having little admixed mineral material.
    1. Brown or yellow strongly acid peat.

As can be seen, this system may be expanded to include more divisions as a study of the organic soils in any area may require.

**The Soil Series.** A soil series, as previously mentioned (p. 43), is usually given the name of a town, river, or other geographical or political feature near which it was first identified or mapped. The series name carries with it a knowledge of all the characteristics involved in differentiating the series. Thus, to one who is familiar with soils the mention of a series name conveys a mental picture of the soil—topography, drainage conditions, character of profile, reaction, color and structure of the horizons, texture of the subsurface horizons, approximate humus content, and something concerning the general nutrient content and productivity for crops commonly grown. It is thus seen that a series name is a highly informative term. However, specific knowledge of the surface soil texture of mineral soils or the degree of decomposition of the surface of organic soils is not conveyed by the series name.

**Soil Type.** A combination of the series name and the texture of the plow soil is necessary to describe mineral soils. This combination of names gives the *soil-type name*. Thus the type name Miami (series) loam (class) brings to the mind of one familiar with the timbered soils of the northeastern Central States a great deal of information concerning a particular soil. In New York Honeoye loam is a very informative name, as is Sassafras sandy loam to people living on the Atlantic Coastal Plain. Cecil sandy loam carries much meaning to persons familiar with

<sup>3</sup> J. O. Veatch, L. C. Wheeting, and A. Bauer, *Soil Survey of Washtenaw County, Michigan*, U. S. Bur. Chem. and Soils, Soil Surveys Series, 1930, No. 21, 47 pp., illus., Washington, D. C.

soils of the Southeastern States. Barnes silt loam brings to the mind of the soil scientist a picture of a widely known soil type in the Central Plains, whereas Mohave sandy loam describes to him a soil of the southwestern desert region. A type name can be considered a short-hand term which briefly and concisely describes one particular soil. Organic soil types are designated muck or peat in addition to the series name; for example, Greenwood peat and Carbondale muck.

**Soil Classification Compared to Botanical Classification.** This fundamental system of classifying soils is similar in many respects to systems used in classifying other natural objects, such as plants and animals. In botany the genus name together with the species name is the common method of technically designating a plant. An analogous binomial method is used in soils, but the above groups are called series and class, respectively. The name Hillsdale sandy loam is comparable to the botanical name *Quercus alba*. Note, however, that English words are used in soils in contrast to the Latin terms of botany. Plants have both a scientific name and a common name. Soils have only one, the soil-type name, which must satisfy both scientific and practical requirements. This name is composed of two parts, the series designation and the class designation. A comparison of the botanical categories and the pedologic categories is shown more fully in Chart 1.

Chart 1. A Comparison of Botanical and Pedologic Classification

| BOTANICAL CLASSIFICATION |                       | PEDOLOGIC CLASSIFICATION |                     |
|--------------------------|-----------------------|--------------------------|---------------------|
| Phylum                   | <i>Spermatophyta</i>  | Order                    | Zonal               |
| Order                    | <i>Angiospermae</i>   | Suborder                 | Forest              |
| Class                    | <i>Dicotyledoneae</i> | Great soil group         | Gray-brown podzolic |
| Family                   | <i>Fagaceae</i>       | Family                   | Miami               |
| Genus                    | <i>Quercus</i>        | Series                   | Hillsdale           |
| Species                  | <i>alba</i>           | Class                    | Sandy loam          |
| Variety                  | .....                 | Phase                    | Hilly               |
| Common name              | White oak             | Hillsdale sandy loam     |                     |

Another difference between the botanical system and the soil system is that the latter is much younger and is therefore still changing. It has developed only since the turn of the century. Not enough is yet known about soils to justify writing a book on their classification similar to Gray's *New Manual of Botany*. However, descriptions of each of the

more than 5,000 soil series now officially recognized in the United States can be obtained from the Division of Soil Survey of the U. S. Department of Agriculture, or from the agricultural experiment stations in the various states.

New soil types are being recognized continually. To get information about these types it is necessary to consult the soil survey reports that will be discussed in a later section. There is still considerable confusion concerning these soil types, however, because of the extreme youth of the science.

**Soil Orders.** The highest category in the classification of soils is the *order*, of which there are three—*zonal*, *azonal*, and *intrazonal*. Zonal soils cover wide areas. Included are those great soil groups, possessing well-developed profiles, that reflect the influence of the active factors of soil genesis, especially climate and vegetation. Topographically, zonal soils are situated on gently undulating, well-drained uplands in parent material not extreme in its texture or chemical composition. Zonal soils are discussed briefly on pages 84–89.

The intrazonal soils have well-developed characteristics that reflect the dominating influence of some local factor, such as the nature of the parent material, topography, or drainage over the normal effect of climate and vegetation.

Azonal soils are without well-developed profile characteristics because some factor of parent material, relief, or age has prevented their development. The relationships of the soil orders are shown in Chart 2 (p. 92). The soil orders are divided into suborders.

**Suborders of Soils.** It will be noticed that suborders of soils are set apart on the basis of the factors which have been most instrumental in determining their characteristics or on the basis of the distinguishing characteristics of the soil group itself. For example, the cold and arid soils are direct products of climatic conditions. Grassland, transitional, and forest soils owe their properties to a combination of the influence of climate and vegetative cover. Excess moisture is responsible primarily for the nature of hydromorphic soils, and the accumulations of chlorides and of calcium salts are the distinguishing characteristics of halomorphic and calcimorphic soils, respectively. The suborders are divided into the great soil groups. See Chart 2.

This soil classification system is the one currently used in the United States. It is really a combination of the genetic system of classification of the Russians and the natural soil classification system developed in the United States. The higher categories from the great soil groups up through the suborders and orders are largely differentiated on a genetic

basis. The lower categories, soil series, types, and phases are based entirely on properties of the soil profiles.

## SOIL MAPPING AND THE SOIL SURVEY

Two subjects closely related to soil classification are soil survey and soil mapping. It was not until a systematic study of the soil resources of the United States was begun in 1898 that the need for a logical, natural system of classification was encountered. And only through the processes of surveying and mapping have the many different soil categories been recognized and correlated. A few questions will assist in fixing in the mind of the student the primary points concerning soil surveying.

### Questions

1. What is a soil survey?
2. What are the steps in making a soil map?
3. What kinds of soil surveys are there?
4. What are the uses of soil survey reports and maps?

**The Soil Survey.** At the turn of the century there was an increasing awareness of the bonds between land and society. In an attempt to find the underlying causes of some agricultural problems and in an effort to build a solid foundation for future research, the U. S. Department of Agriculture in cooperation with the various state experiment stations began at that time a systematic investigation of our soil resources. This investigation assumed the form of a national inventory and survey.

At the present time soil surveying is the process of studying and mapping the earth's surface in terms of units called soil types. A soil survey report thus consists of two parts: (1) the soil map, which is accompanied by (2) a description of the area shown on the map.

**Soil Mapping.** Surveys and maps are made for county areas. The actual process of mapping consists in walking over the land at regular intervals, generally about 80 rods, and taking notes by means of a "legend" and "field sheet" on soil differences and all related surface features, such as slope gradients, evidence of erosion, land use, vegetative cover, and cultural features. The field sheets are sometimes geological survey base maps, sometimes plane tabled by the soil surveyor himself, but in recent years contact prints of aerial photographs have become increasingly popular. Boundaries are drawn directly on the field

sheets, representing in most places changes from one soil type to another.

Soil survey activities within a state are usually under the direction of the state experiment station. This organization and other state and federal agencies share the expenses of mapping with the Soil Survey Division of the U. S. Department of Agriculture, which inspects the work from time to time in order not only to maintain a certain working standard but also to correlate the mapping units in all sections of the country and to confirm the recognition of new soil types, phases, and series. After the field work has been completed, the data are assembled, printed, and made available to the public by the U. S. Department of Agriculture.

There are over 2,850,000 square miles in the United States that are considered to possess reasonable possibilities for the production of crops, grasses, and trees. Of this area, approximately 942,500 square miles have been mapped, although some 325,000 square miles need to be resurveyed because of advances in knowledge since the work was done. There are, then, somewhat more than 1,000,000 square miles which should be surveyed in detail, with 600,000 square miles for which semi-detailed maps are needed and an equal area to be surveyed on a reconnaissance basis. These figures include some work needed in Alaska and Puerto Rico. Some 1,600 survey reports, including over 1,800 maps mostly on a county basis, have already been published.

**Types of Soil Surveys.** Productive agricultural land is studied and mapped in detail. A detailed map is one in which the scale is 1 or more inches to the mile. Four inches is the scale usually used at present, although occasionally an 8- or even a 12-inch scale is employed. The purpose is to supply information necessary in planning land-use operations on individual farms and fields. In addition to soil-type boundaries, these maps show the extent of removal of the surface soil by erosion, location of gullies, gradient and direction of slopes, and other details needed in planning a soil-conserving, practical management plan. This kind of work is expensive, costing from \$75 to \$150 for each square mile, that is, from 12 to 24 cents per acre. Large expanses of land are too poor, however, to justify this cost. These areas are mapped on small scales, and the resulting study, called a reconnaissance survey, shows areas and regions that are dominated by a single soil type, in contrast to the detailed map where all types and phases are separated.

In recent years still another kind of soil mapping has been employed, particularly in land of low agricultural value. Called *land-type mapping*, it consists in studying *soil-type associations* in contrast to merely separating areas in which a single type is dominant, as in reconnaissance

surveying. Land-type maps are useful in studying extensive land-use problems, and they possess the desirable characteristics of being made quickly at a comparatively low cost.

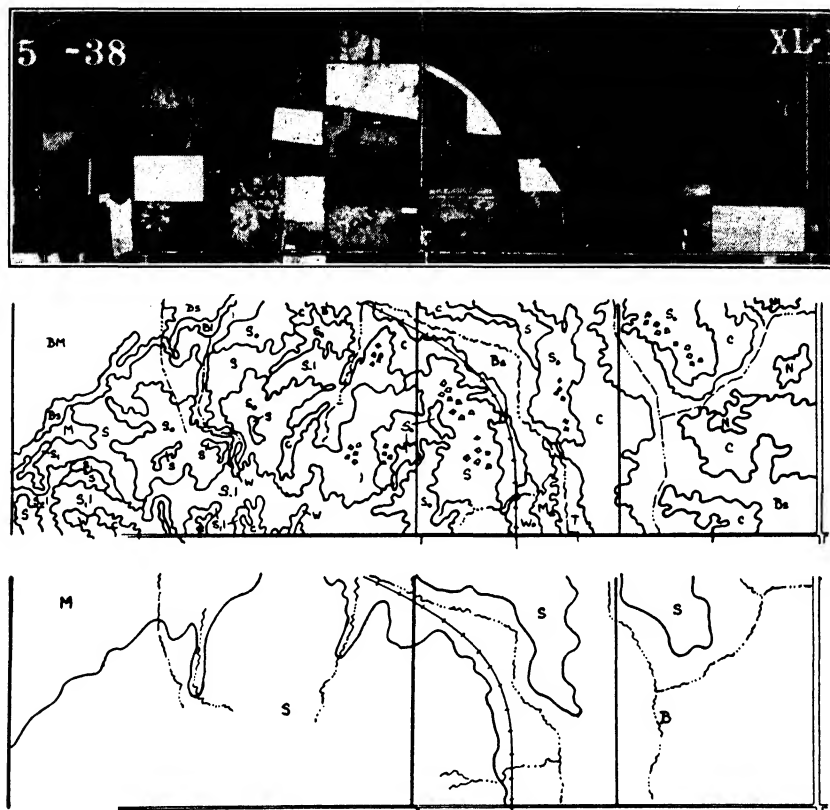


FIG. 21. Top: Aerial photograph of good farming land in glaciated area. Such photographs are used extensively for a base map in making soil surveys. Center: A detailed soil survey map of the same area. The symbols indicate soil types. Bottom: A land-type map of the area in which soil types of similar characteristics occurring in close association are mapped as a unit.

**Purpose and Value of Soil Surveys.** The general purpose of a soil survey is to obtain an inventory of the soil resources of the country at large, of individual states, and more locally of counties and communities. More specifically, a soil survey gives information concerning soil quality, including topography, natural drainage, natural fertility, susceptibility to erosion, general crop adaptation, need for lime, and so forth, of any

given community. Often this information can be applied to individual farms.

The soil survey is used by federal and state agencies, including colleges and universities, to obtain dependable information concerning soil conditions in given areas. Life insurance companies, banks, and other money-lending agencies use soil surveys in determining security for

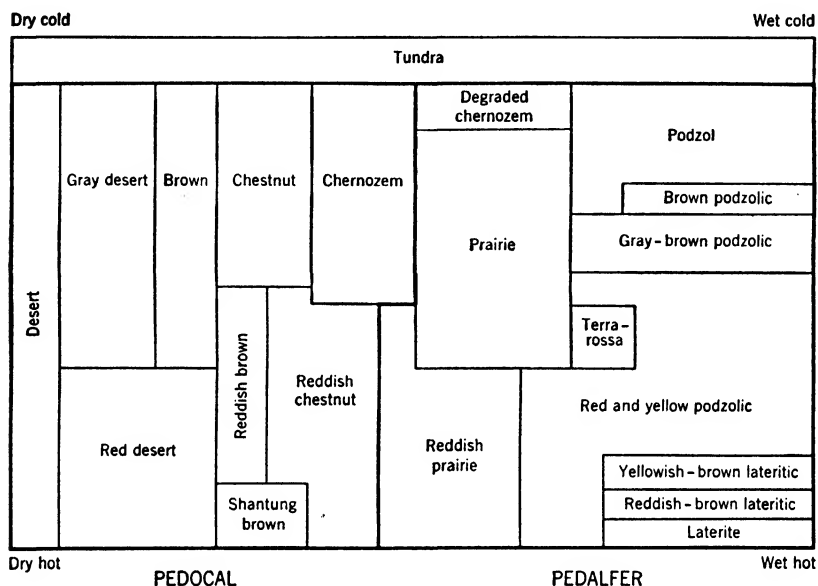


FIG. 22. A diagrammatic visualization of the relation between climatic conditions and the occurrence of the great soil groups. The relative size and position of each area indicates the extent and general geographic location of the soil group represented.

loans. Real estate companies and individuals interested in buying or selling land make extensive use of soil surveys. They are also used by highway and drainage engineers, by various kinds of manufacturers in selecting suitable locations for factories, and by merchandising and advertising companies in selecting areas for intensive campaigns. County agricultural agents and other extension workers find soil surveys helpful in their work. And, last but by no means least, the farmers themselves are making increasing use of soil survey maps and reports in planning their management programs and in interpreting modern agricultural research in terms of their own farm conditions.



## GENERAL DISTRIBUTION OF THE GREAT SOIL GROUPS

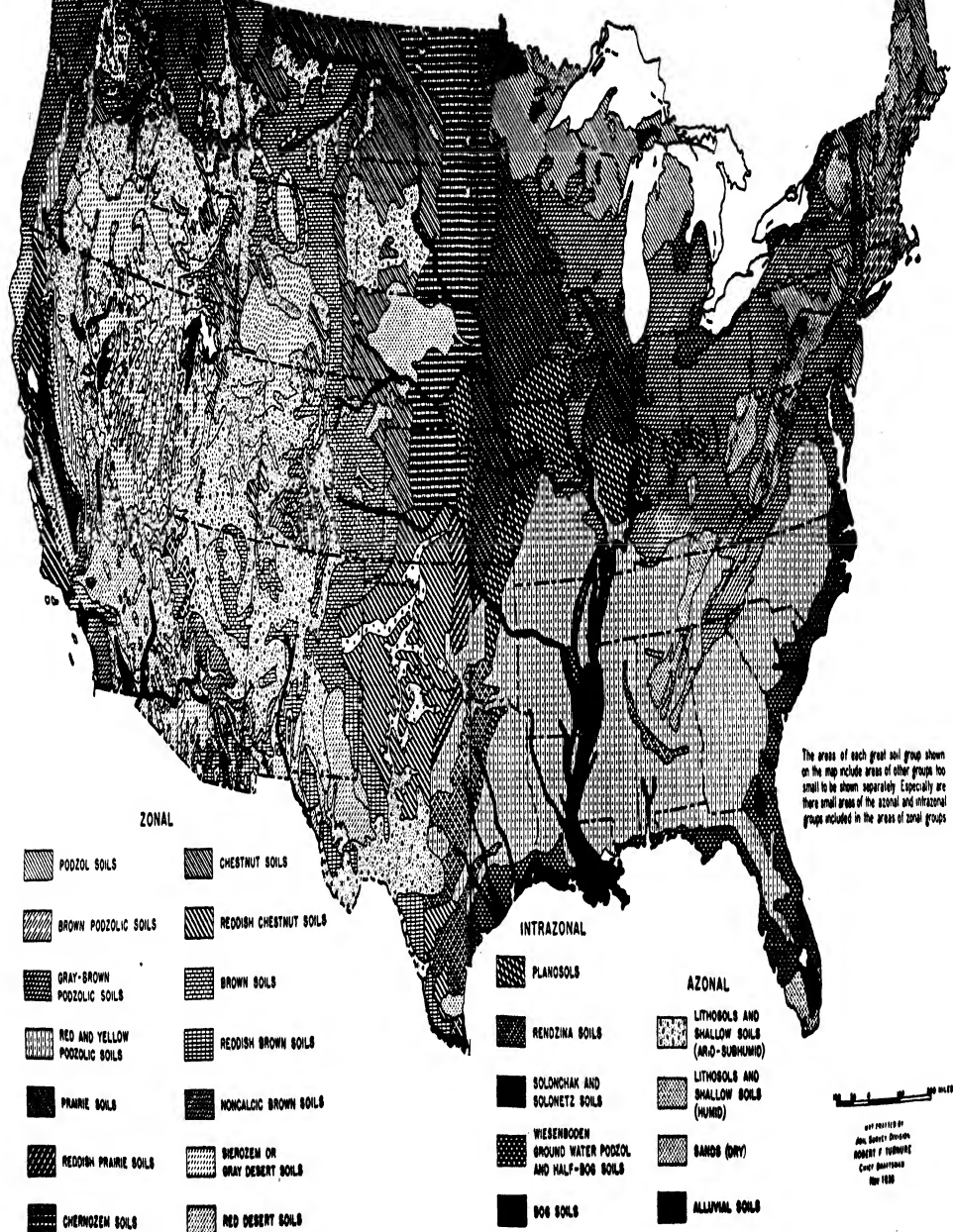


FIG. 23. General distribution of the great soil groups in the United States. [Courtesy of Division of Soil Survey, U.S.D.A.]



## SOIL GROUPS IN RELATION TO CLIMATIC CONDITIONS

The information available from soil maps and soil-survey reports makes possible broad geographic studies on the relationship between soils and their environmental conditions. Of these relationships none is more evident or of greater interest than that which exists between the great soil groups and climate.

The rate of activity of the various forces of soil formation is greatly influenced by climatic conditions, as has been previously pointed out. Furthermore, the disposition of certain of the products of mineral decomposition is determined by the distribution, the quantity, and the nature of the precipitation. In view of these facts, it is inevitable that different climates must be related to the development of different profile characteristics. A study of Fig. 22 shows that there is a broad or general grouping of soils according to the climatic zones in which they were formed. The following questions emphasize important factors in these relationships. Limitation of space makes it necessary to confine the discussion to a few of the zonal soils.

### Questions

1. May soils be divided into groups on the basis of the leaching of salts from the surface layers?
2. In what soil groups was there a movement downward of iron and aluminum?
3. What soil groups contain an accumulation of salts?
4. What kinds of soils are found in the tropics?
5. How are Tundra soils developed?

**Leached and Unleached Soils.** Soils developed in a region where precipitation is sufficient to remove from the soil profile the very soluble salts of potassium and sodium and the less soluble carbonates of calcium and magnesium differ in many respects from soils developed in a climatic zone where precipitation is insufficient to remove these compounds. A line drawn north and south across the United States near Lincoln, Nebraska, roughly marks the division between these two climatic zones (Fig. 23).

**Soils in which Iron and Aluminum Have Leached.** Not only are soluble salts leached readily from soils east of this dividing line, but also a downward movement of iron and aluminum compounds takes place. Suspended organic matter also moves downward. This leaching, which commonly results in a whitening of the soil layer immediately below the surface organic matter and the accumulation of iron and

aluminum compounds in the underlying brown, yellowish brown, yellow, or red *B* horizon, is known as podzolization. The strongly leached part of the soil (*A* horizon) is known as the *horizon of eluviation*. The part in which compounds of iron, aluminum, and other elements are concentrated is known as the *horizon of illuviation*. See Fig. 11 on page 37.

The soils in this climatic zone are given a general or group name based on this tendency for iron and aluminum to segregate within the profile. The first syllable of the Greek word *pedon*, meaning ground, is coupled with abbreviations of the Latin words *alumen* (aluminum) and *ferrum* (iron) to form the name *pedalfers*.

*Characteristics of Prairie Soils.* Precipitation increases as one moves eastward from Lincoln, Nebraska. Precipitation is accompanied by a corresponding increase in height of the prairie grasses. In what is now the heart of the Corn Belt, the native grasses originally stood as high as a man's head. A deepening of the horizon of humus accumulation follows to some extent the increase in height of the grasses until a depth of 20 or more inches is reached. Accumulation of iron, aluminum, and clay in the illuviated horizon increases with increasing precipitation, and the horizon of eluviation becomes more apparent. Consequently, these soils are pedalfers. To a considerable extent the prairie soils developed in loessial and glacial materials of Wisconsin age, and the topography varies from level to rolling with predominantly long and gentle, rather than steep, slopes. The term "Brunigra" (brownish black) has been proposed to replace Prairie as the name of this great soil group.

*Gray-Brown Podzolic Soils.* Eastward from the prairies to the Atlantic Coast the soils were timbered. In the area bounded on the north by a line running east and west in the vicinity of Milwaukee, Wisconsin, and on the south by a line through Cairo, Illinois, are found the Gray-brown Podzolic soils. In these soils the horizon of humus accumulation (*A*<sub>1</sub>) is comparatively thin, ranging from 2 to 6 inches in thickness. The light-colored eluviated horizon (*A*<sub>2</sub>) is well-developed, varying from grayish to yellowish brown. The horizon of illuviation is conspicuous because its color, varying from grayish to reddish brown, is generally darker than the color of either the parent material or the overlying leached horizon.

*Podzols.* North of the region of gray-brown podzolic soils, in a climate of lower temperatures and higher humidity (due to low evaporation), are developed the *Podzols*. Here the humus horizon (*A*<sub>0</sub>) is of moderate thickness, the bleached, eluviated horizon (*A*<sub>2</sub>) is strongly developed, and the horizon of illuviation is conspicuous. The latter is

usually brown to coffee-brown in color and is often characterized by a slightly cemented or indurated structure. Organic compounds together with iron oxides serve as cementing agents. The podzol profile represents the highest development of the podzolization process in soil formation.

*Red and Yellow Soils.* South of the region of gray-brown forest soils, high temperatures coupled with heavy rainfall accelerate the rate of mineral decay. The leaching of soluble products as well as the decay of organic matter is also increased. Soils developed in this climate are therefore characterized by a low accumulation of surface organic matter, a deep horizon of eluviation, and a deep, thick illuviated horizon in which the high rate of oxidation and hydration of iron produces bright red and yellow colors. The southern boundary of the Red-and-Yellow-soil area, where there is evidence in many places that the present soils were derived from old profiles developed under a more tropical climate than the existing one, merges into the lateritic zone.

**Soils Containing Salt Accumulations.** Soils in arid and semi-arid regions are characterized by an accumulation of calcium carbonate or by a mixture of  $\text{CaCO}_3$  and  $\text{MgCO}_3$  at some position in the profile. In recognition of this characteristic the Greek word *pedon* is combined with an abbreviation of the Latin word *calcis* or *calx* (lime) to produce the name *Pedocal*.

In this group of soils the colloidal iron and aluminum oxides do not disperse and move downward in the profile because the soils retain an abundance of active bases. The neutral to alkaline reaction prevents their dispersion.

*Chernozems.* In a wide north-south belt occupying the eastern part of the Great Plains of central United States is developed a group of soils containing little of the soluble salts of potassium and sodium but a striking accumulation of calcium salts occurring at a depth of approximately 24 to 36 inches. The natural vegetation of this region is composed of mixed prairie grasses, under which a relatively thick, granular, humus-mineral layer has formed. In places as much as 15 to 20 inches of dark surface material develop; consequently these soils are known as "black soils." They bear the name of *Chernozem*, which was given to them when they were first classified on the steppes of Russia.

*Chestnut Soils.* As one travels westward from the chernozem belt into zones of less rainfall, the accumulation of calcium salts occurs nearer the surface (14 to 24 inches) and salts of sodium and potassium are present in increasing quantities. The vegetation becomes more sparse and shorter in stature. The very dark color of the surface soil

horizon gives place to a brown, and this layer becomes thinner. This belt, lying just west of the chernozems, constitutes the *Chestnut soils*.

*Brown Soils.* West of the chestnut soils is an area of light brown or grayish brown soils. The surface horizons are lower in humus content and thinner. They support a still shorter and sparser grass vegetation intermixed with some shrubs and are richer in soluble salt content.

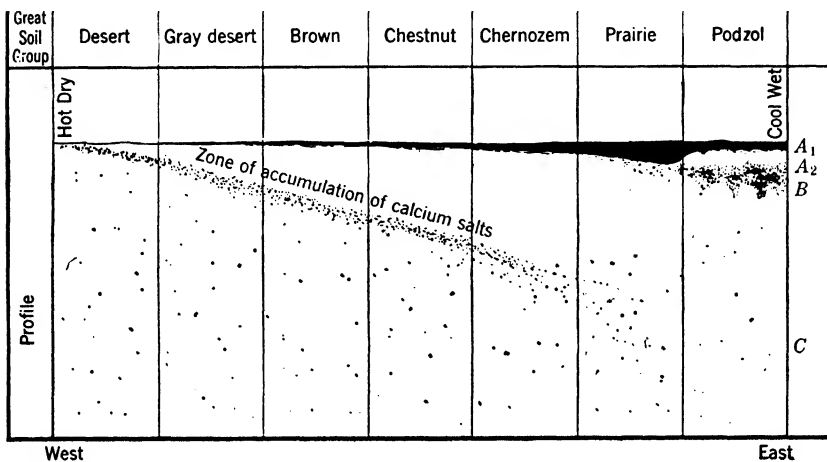


FIG. 24. Generalized profiles of the zonal soil groups in central and western United States. Note how the zone of accumulation of calcium salts increases in depth and diminishes with increase in precipitation. The thickness of the humus-rich layer also increases with increased precipitation until the grass vegetation of the prairies gives place to the forests of the Podzolic region. The development of a clay-rich *B* horizon is prominent in the Podzolic soils but disappears as one proceeds westward in the prairies.

A horizon of accumulation of calcium carbonate occurs at a depth of 12 to 15 inches or less.

*Desert Soils.* Under desert conditions the humus content of soil becomes so low that the brown of semi-arid regions is replaced by gray, which is intensified by a surface accumulation of salts. Usually a few inches of soil rest on the calcareous parent material. Shrubs replace grass, and cacti appear in the warmer areas. Concentration of soluble salts is sufficient in some areas to produce "alkali" spots, which support only alkali-resistant plants.

*Soils of the Tropics. Latosols.* There is a great diversity of climatic conditions in the tropical zone. There has been an inclination to think of the "tropics" as an area of high rainfall, high temperature, and

dense forest growth. Although there are extensive areas of this nature there are also deserts, swamps, and snow-capped mountains. Soil materials also vary greatly. In consequence, there is much variation in tropical soils. One formation of the tropics which has attracted much attention is composed largely of the more or less hydrated oxides of iron and aluminum, and called a "laterite." The term laterite came to be used as a name of a great soil group, with the inference that all tropical soils are of a somewhat similar nature. This classification is now being reviewed. Further study of tropical soils has led Dr. C. E. Kellogg to suggest that the term laterite be applied only to "sesquioxide-rich, highly weathered clayey material that changes irreversibly to concretions, hard-pans, or crusts when dehydrated, and for the hardened relics of such material more or less mixed with entrapped quartz and other diluents. It is further suggested that the term 'latosol' be defined and used as the name of the suborder of zonal soils of the tropics." Soils of the tropical zone need much more study; their great diversity should be kept in mind.

**Tundra Soils.** Tundra soils are composed of a mat of living and dead grasses, mosses, sedges, and lichens. Little is known of the soil material underlying this mat, and apparently it has little effect on the accumulation of plant growth on the surface. The tundra occupies large land areas where the subsoil remains frozen throughout the year: those areas bordering the Bering Sea and the Arctic Ocean. The native vegetation varies, but on the average it consists of 30 per cent lichens, 25 per cent sedges, 25 per cent shrubs, and 20 per cent grasses, weeds, and mosses. The tundra is capable of furnishing grazing for large numbers of reindeer, and some movement to utilize it for this purpose has been started.

#### AGE, RELIEF, AND PARENT MATERIAL IN RELATION TO SOIL GROUPS

In the preceding sections the soil profiles in some of the great soil regions have been discussed briefly. It must be remembered, however, that by no means all soils in a zone have profile characteristics that are identical with, or closely approach, those of the zonal profiles. Unfavorable drainage conditions, topography, time during which soil-developing processes have been active, and the great resistance of some parent material to weathering are among the factors which account for the lack of uniformity within a region. In discussing soil areas it should be also recognized that usually the properties of one soil change gradually as neighboring soils are approached; just as there is no sharp line

of division between soil types, there is no sharp line of division between great soil groups. Figures 12, 13, and 14 illustrate the influence of age, topography, and parent material, respectively, on soil-profile development.

### SOIL GROUPS IN RELATION TO VEGETATIVE COVER

Natural vegetative cover may be divided into the two general classes of trees and grass, and the soils supporting them are termed forest soils and grassland soils, respectively. There are several characteristics in soils developed in association with grass which are of considerable agricultural significance. Furthermore, there are two types of forest, coniferous and deciduous, and two types of prairie grasses, the tall and the short. The different effect of each kind of vegetation on the soil supporting it is brought out in the answers to the questions listed.

#### Questions

1. Why is there a difference in humus content of prairie and forest soils?
2. Are equal quantities of nutrients returned to the soil annually by grass and trees?
3. Is action of microorganisms the same in prairie and timbered soils?
4. Does leaching proceed as rapidly under grass as under trees?

**Humus Accumulation.** When grassland and timbered soils are brought under cultivation, one of the first differences between them to attract the attention of a casual observer is the contrast in color of the surface layer. Under similar drainage conditions grassland soils are much darker, owing usually to a higher humus content but sometimes to a difference in the quantity of calcium combined with the humus. The humus-rich horizon in these soils is also found to extend to a much greater depth. A partial explanation of these differences is found in the nature of the root systems of the two types of plants.

The fibrous-root system of grass completely fills the upper part of the soil with minute rootlets, which upon decaying leave the resulting organic matter thoroughly distributed within the surface layers. Tree roots, on the other hand, are large, with quantities of fine rootlets occurring very near the surface instead of being scattered in depth. In fact, many of them extend up into the more decayed sections of the litter. Only a thin layer of forest soil, therefore, is enriched in organic matter through rootlet decay. As the larger roots rot, the resulting organic matter is left in channels where it has no chance to be widely distributed throughout the soil.

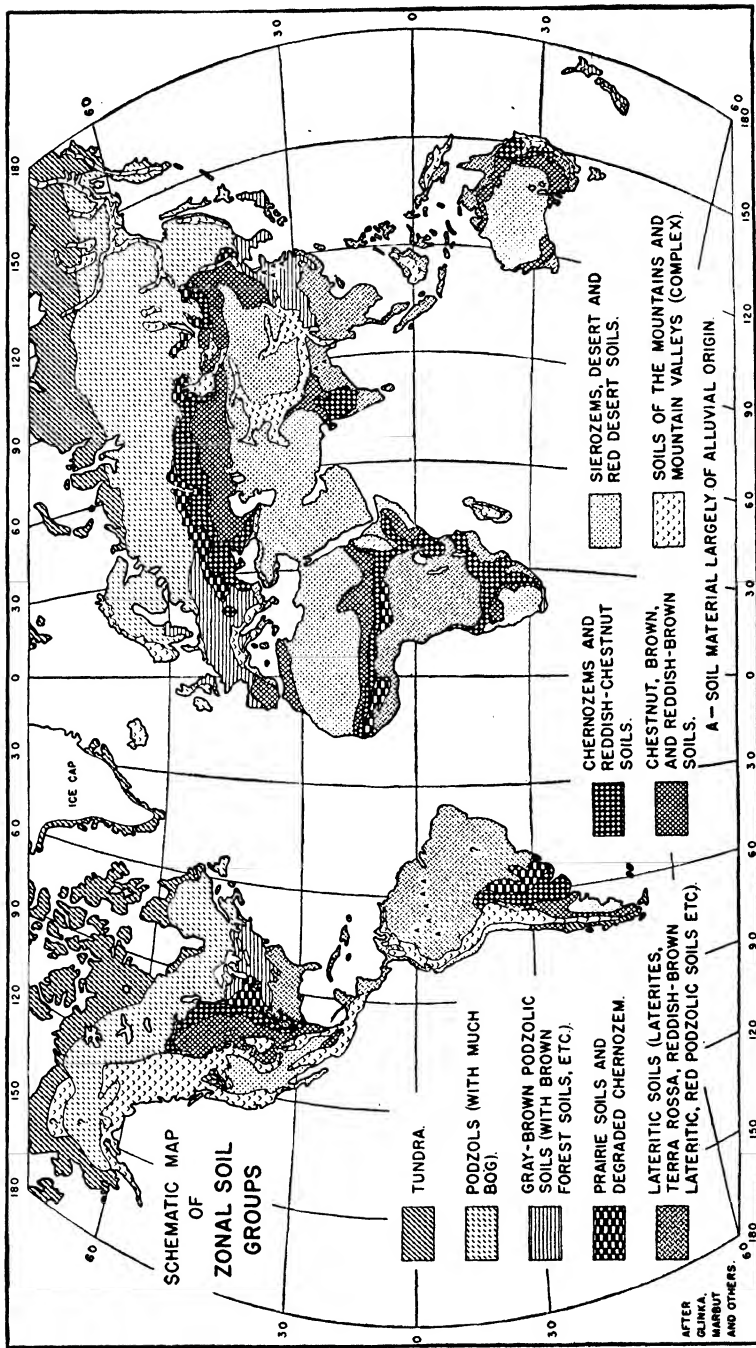


FIG. 25. A generalized map showing the approximate distribution of the zonal soil groups throughout the world. [From *The Soils that Support Us*, by C. E. Kellogg, The Macmillan Company, 1941.] See page 88 for a more recent discussion of lateritic and tropical soils.

## Chart 2. A Classification of the Great Soil Groups \*

| Order            | Suborder   | Great Soil Groups  |
|------------------|--|--|
| Zonal soils      | 1. Soils of the cold zone  | Tundra soils   |
|                  | 2. Light-colored soils of arid regions   | Desert soils<br>Red desert soils<br>Sierozem<br>Brown soils<br>Reddish brown soils   |
|                  | 3. Dark-colored soils of semi-arid, subhumid, and humid grasslands                                 | Chestnut soils<br>Reddish Chestnut soils<br>Chernozem soils<br>Prairie soils<br>Reddish Prairie soils  |
|                  | 4. Soils of the forest-grassland transition  | Degraded Chernozem<br>Non-calcic Brown or Shantung Brown soils   |
|                  | 5. Light-colored podzolized soils of the timbered regions  | Podzol soils<br>Gray wooded or Gray Podzolic soils †<br>Brown Podzolic soils<br>Gray-brown Podzolic soils<br>Red-yellow Podzolic soils †   |
|                  | 6. Lateritic soils of forested warm-temperate and tropical regions                                 | Reddish brown Lateritic soils †<br>Yellowish brown Lateritic soils<br>Laterite soils †   |
| Intrazonal soils | 1. Halomorphic (saline and alkali) soils of imperfectly drained arid regions and littoral deposits | Solonchak or<br>Saline soils<br>Solonetz soils<br>Sloth soils  |
|                  | 2. Hydromorphic soils of marshes, swamps, seep areas, and flats                                    | Humic-gley soils † (includes <i>Wiesenboden</i> )<br>Alpine meadow soils<br>Bog soils<br>Half-bog soils<br>Low humic-gley † soils<br>Planosols<br>Ground water Podzol soils<br>Ground water Laterite soils |
|                  | 3. Calcimorphic soils  | Brown forest soils ( <i>Braunerde</i> )<br>Rendzina soils  |
| Azonal soils     |  | Lithosols<br>Regosols (includes dry sands)<br>Alluvial soils   |

\* Prepared by James Thorp and Guy D. Smith, published in *Soil Science*, Vol. 67, No. 2, p. 118. Used through courtesy of Williams and Wilkins Co.

† New or recently modified great soil groups.

The aerial portions of plants differ widely in their method of contributing organic matter to the soil. Grass falls more or less evenly on the ground, whereupon the dead stems and leaves may be dragged under the surface by worms and insects. The products of decay are also carried into the soil by water. Only a small part of the annual growth of trees, however, is returned to the soil surface in twigs and

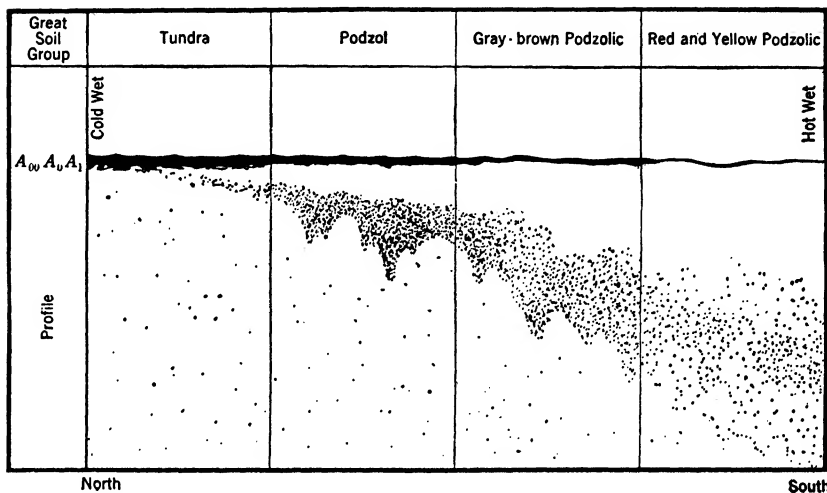


FIG. 26. Generalized profiles of zonal soil groups in eastern North America. Note how the accumulation of organic matter decreases with increase in temperature. The mineral material under the surface mat of organic matter in the Tundra is very slightly altered. In the Podzol zone eluviation is intense, and a more or less indurated *B* horizon containing much humus and iron oxide is developed at shallow depths. In warmer climates eluviation extends to greater depths; the *B* horizon is thicker and is richer in clay with less iron and humus than in the Podzol. The reds and yellows of the *B* horizon in the Red and Yellow Podzolic soils are due to the state of oxidation and hydration rather than to an excessive quantity of iron.

leaves. They contribute to the humus content of the soil in the same manner as grass tops if worm and insect action are equal, but frequently they are not. When large branches and tree trunks fall, the products of decay are deposited on limited areas instead of uniformly over the entire surface, as with grass tops. When forest land is being cleared for cultivation, part of the leaf litter, dead logs, and branches are burned and hence never contribute their quota of humus to the soil.

**Difference in Nutrient Cycle.** Because of the completeness with which grass roots fill the soil and the annual habit of growth, large

quantities of minerals are carried up to the leaves and stems and returned to the surface soil each year. This return of minerals to the surface layers results in a larger supply of available nutrients for the succeeding growth than there would be under timbered conditions, and it also

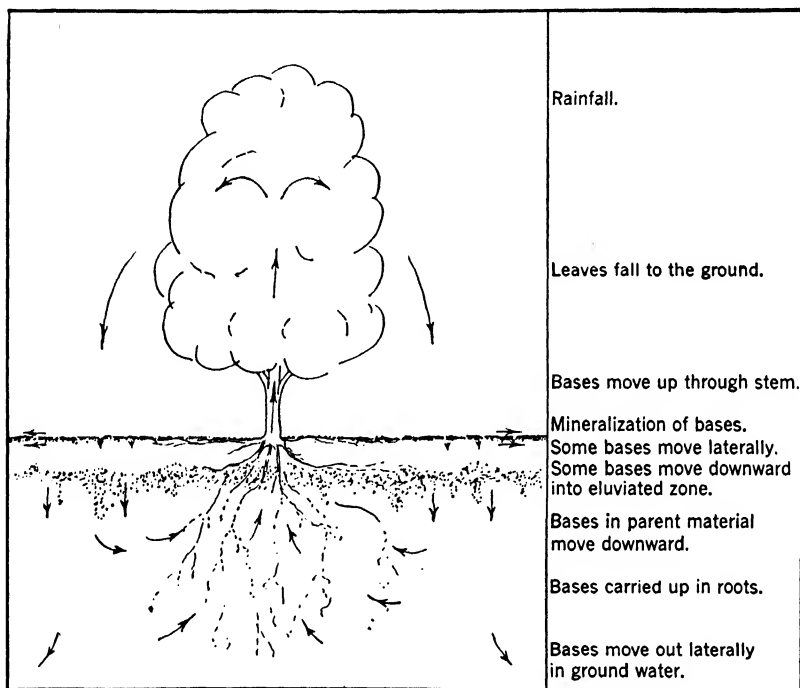


FIG. 27. The base movement in a forest soil. The base cycle has a strong influence on the properties of the soil developed under different types of vegetative cover. On the prairies all the leaves and stems of the grass with their content of bases are returned to the soil surface annually. In the forest a decidedly different cycle occurs since much of the nutrient supply taken up by the roots is stored in the tree trunks and branches.

delays the tendency of the soil to become acid. A grass cover also decreases the leaching away of soluble nitrogen compounds and hence leads to a further accumulation of humus because of the vital part played by nitrogen in humus formation (see Chapter 10).

Much of the mineral and nitrogenous plant-nutrient content of leaves of trees is translocated to the woody parts of the plant before the leaves fall, and hence there is not the large annual return of these materials to the surface soil under a forest cover than there is under a grass cover.

**Action of Microorganisms.** In grassland soils both bacteria and fungi are active. On the other hand, under forest cover fungous action is thought to exceed bacterial action. This is especially true under coniferous forests. The products of the two types of decomposition are somewhat different, those resulting from the action of fungi being more acid in nature. This difference in the type of microorganic population may account in part for the difference in humus content of soils under the two types of cover and the more rapid leaching under forest conditions.

**Eluviation Slower in Grassland Soils.** Because of the nature of the root system of grass which fills the soil completely, much soluble nutrient material is absorbed which would otherwise leach away. The high content of colloidal organic matter (humus) also provides for the retention or "fixation" of large quantities of nutrients. These factors, together with the large annual return of bases to the surface soil, are in part responsible for the slowness of development of a bleached eluviated horizon in grassland soils. Such a horizon is found much more highly developed in timberland adjoining a piece of grassland than in the grassland soil.

#### SOIL GROUPS IN RELATION TO POPULATION DENSITY AND PRODUCTION OF AGRICULTURAL PRODUCTS

Large cities generally develop in regions which produce large quantities of agricultural products. Some exceptions are due to the necessity of utilizing harbors for water transportation and of operating mines in non-agricultural areas. Not only do agricultural workers produce many of the articles required by factory workers, but also they are themselves large consumers of the products of the factories. Moreover, a large number of persons are engaged in processing and marketing products of the soil. The mutual interests of the city and farm worker are many, and the common recognition of this fact will benefit everyone. The following questions will serve as guides in acquiring some knowledge of the agricultural products of each of the great soil groups and of the nearness of these areas to population centers where such products are consumed.

#### Questions

1. What type of farming is practiced on the Chernozems?
2. How are the Chestnut soils farmed?
3. Are the Brown soils used for cultivated crops?

4. What agricultural use is made of the Sierozem Desert-soil areas?
5. What farming conditions prevail on the Prairie soils?
6. How are the population density and agricultural development of the Gray-brown Podzolic area accounted for?
7. What type of agriculture do the Podzols support?
8. Why are the Red and Yellow soils farmed so extensively?
9. How are organic soils farmed?

**Chernozem Agriculture.** Population is sparse in the Chernozem belt because farming must be extensive rather than intensive, owing to limited rainfall. Chernozems are noted for their production of wheat not only in the United States but also in Russia and wherever climatic conditions have produced such soils. Other drought-avoiding and drought-resistant crops, such as varieties of the sorghums, are also grown. Corn and oats are produced to a limited extent, as well as considerable cotton in the South. Livestock raising is an important part of the farm program.

The soils are well supplied with mineral nutrients and nitrogen and are highly productive when supplied with moisture. The annual precipitation averages 20 to 26 inches. A mixture of tall and mixed grass is the prevailing native vegetation.

**Agriculture of the Chestnut Soils.** These soils constitute a considerable part of the spring wheat belt and also produce some sorghum crops for grain and forage and a little corn. Stock raising assumes a much greater relative importance than on the Chernozems, and extensive herds of cattle graze on the nutritious native grasses. The population is more widely scattered, for pastures must be large because of their relatively low carrying capacity. Ample water would make these soils highly productive, but the meager rainfall (15 to 22 inches) materially limits crop production.

**Brown-Soil Farming.** These soils cover a vast area in the western part of the Great Plains with smaller areas in the intermountain region of the far West. The climate is semi-arid with temperate to cool temperatures. The native vegetation consists of short grasses, bunch grasses, and shrubs. The proportion of the land planted to crop varies from virtually none in some areas to most of it in others. Dry farming is practiced extensively, and irrigation is utilized to a certain extent. A great variety of crops is grown, especially when irrigation is used. Wheat is grown extensively, as are also the grain sorghums, corn and alfalfa. Fruits, vegetables, and sugar beets occupy smaller acreages. Grazing is a major form of land use, and in some sections most of the land is devoted to cattle raising. Grazing is extensive, and large shipments of cattle move each year from this and the Chestnut-soil zone

to the feed yards of the Corn Belt. Dried buffalo and grama grasses afford winter grazing for cattle.

**Land Use in Sierozem Desert-Soil Areas.** These soils support primarily a shrub vegetation with bunch grasses of varying thickness and a growth of short-lived annuals in the spring or after rains. Without irrigation the land is suited only for grazing and has a very low carrying capacity. In some sections sheep raising is the main enterprise. Where irrigation has been developed, a variety of crops is produced including alfalfa, sugar beets, fruits, grain crops, and vegetables.

**Agriculture of the Prairie (Brunigra) Soil Areas.** The tall-grass prairies lying east of the Chernozems are the scene of extensive cereal and hay crop production and of intensive livestock feeding. Corn, oats, wheat, and soybeans are the principal grain crops. The clovers, alfalfa, and timothy, either alone or in some combination, are used for hay and pasture. Farms are comparatively large and are highly mechanized. Hog raising and cattle fattening are important enterprises. Dairying has developed extensively in the vicinity of larger towns and cities. Farming population is comparatively dense, and many moderate-sized cities have grown up to provide market places and processing facilities for farm products and to make implements and other manufactured goods for the farmer.

Annual precipitation ranges from 28 to 45 inches with a goodly proportion falling during the growing season. A midsummer drought is characteristic of the region, but nevertheless it contains a high percentage of the best farm land of the United States.

**Farming in the Gray-Brown Podzolic Zone.** Farming in this zone is more intensive, with somewhat smaller farms, than in the Prairie region. A larger variety of crops is grown, with less emphasis on corn and more attention to poultry raising, dairying, and truck crop production. Cattle and lambs are fed, and a large number of hogs are raised. The population density is greater, both urban and rural, than in the prairie regions, and a larger number of factories are found in both cities and small towns.

Precipitation ranges from 30 to 50 inches, and, although summer droughts are frequent, moisture relations are on the whole satisfactory. The growing season is long, and summer temperatures are not excessively high except in the southern part of the zone. Soils are of moderate fertility, having suffered considerable leaching and varying losses from erosion. The humus content is medium. It is the combination of favorable climatic conditions and moderate to good soil fertility, permitting a highly diversified agriculture, that has made the Gray-

brown Podzolic areas of the world the scenes of a highly developed agriculture and the location of population centers.

**Farming the Podzols.** Long winters and short, cool summers mark the Podzol zone. Soil moisture is plentiful because of low evaporation. Soil fertility varies from low to high. Hay, potatoes, and roots are the main crops, with a limited production of grain. Because of the abundant forage, dairying is common. Butter and cheese are the main products because they are easily shipped considerable distances to market, which fact is important because the Podzols support only a small local population. Cities in this soil zone owe their existence to lumbering and mining rather than to local agriculture.

**Crop Production on the Red and Yellow Soils.** The combination of ample precipitation, comparatively high temperatures, and long growing season, all of which permit the growing of many different crops, has resulted in a high degree of agricultural development in the Red and Yellow Podzolic regions which support a dense population despite the mediocre productivity of the soils themselves. Farms are usually small to medium-sized. An annual precipitation, varying from 40 to 60 inches, has caused severe leaching of the soil and the development of an acid reaction in addition to much soil deterioration through erosion losses. Rapid decay of organic matter results in only a small humus accumulation. These factors, coupled with the high value of many of the special crops grown, have led to the use of large quantities of commercial fertilizers.

Livestock farming is on the increase, but it has not yet developed to the extent desirable, and more diversification in the cropping system is advisable. Aside from cotton, many special crops, such as tobacco, peanuts, vegetables, nuts, citrus fruits, and small fruits, are produced. There are many wood lots and forested areas.

**Farming Organic Soils.** When organic soils are drained in such a way as to remove excess water rapidly but yet maintain the water level at a relatively shallow depth, they may be used for very intensive types of crop production. In the northern states these soils are used for the production of onions, celery, mint, potatoes, cabbage, cranberries, carrots, and other root crops. Corn is produced to some extent, and considerable areas are used as pasture. Late spring or summer and early-fall frosts are the greatest hazard to crop growth. A great variety of special crops is grown on the organic soils of the South and East. Special methods of tillage, coupled with careful and copious fertilization, are required to bring these soils to their highest state of productivity.

# 5

## Chemical Properties of Soils

The chemical composition of soils as expressed in terms of the percentage of different elements present has received much study. However, as knowledge of the interrelationships of plants and soils has accumulated, less emphasis has been placed on the total percentage of elements in the soil and more attention given to the combinations or compounds in which the elements occur. That is, the concept of availability of nutrients to plants has been given more consideration. This has led to a study of ionic exchange and the composition and structure of the exchange complex. In fact, a realization of the controlling role played by the colloidal fraction of the soil in soil productivity has resulted in an intensive study of colloidal materials and the laws governing their activities. The following objectives have been kept in mind in a brief discussion of the chemical properties of soils.

### **Objectives**

- A. The chemical composition of soils.
- B. The colloidal fraction of soils.
- C. Other chemical properties of soils.

### **THE CHEMICAL COMPOSITION OF SOILS**

The solid crust of the earth or lithosphere contains some of virtually all the 95 or more chemical elements which have been discovered. An analysis of this portion of the earth, however, shows that only a few elements make up the major portion of it. Also, as a soil is developed by the processes of weathering, marked losses occur in the quantities of some of these elements and consequent increases in the quantities of others. The accompanying questions refer to some of the pertinent differences in the composition of different soils.

**Questions**

1. What is the composition of representative soil types?
2. Is there a difference in the chemical composition of the soil separates and classes?
3. Do surface soils differ in composition from their parent material?
4. Is the composition of a soil a key to its fertility?

**Composition of Soil Types.** In discussing the chemical composition of soils reference is made to the composition of the surface layer to the depth of plowing unless otherwise stated. This is the portion of the soil from which plants usually obtain a considerable portion of their nutrients and which man alters physically and chemically through tillage operations and additions of lime, fertilizers, crop residues, and green crops. The amounts of the different constituents present in the soil may be expressed as percentage or in pounds per acre calculated on the basis of 2,000,000 pounds of soil, which approximates the weight of a mineral soil to a depth of 6½ to 7 inches over an acre. The composition may be expressed in terms of the chemical elements, as Ca, P, K, or as oxides of the elements, as CaO, P<sub>2</sub>O<sub>5</sub>, K<sub>2</sub>O. In studying tables of data one must observe carefully the terms in which the analyses are expressed.

When consideration is given to the fact that soils are developed from widely different materials such as alluvial deposits along streams, wind-laid silt and clay or sand, glacial drift, volcanic ash, hard rocks like granite and limestone, and under widely varying climatic conditions, under grass, tree, or shrub cover, and that soils are of vastly different ages, it is to be expected that they will vary greatly in chemical composition. Table 2 gives a partial chemical analysis of a few soil types from different sections of the United States.

In these and similar data a few points are outstanding. (1) SiO<sub>2</sub> constitutes a high percentage of all the soils. This would not be true of organic soils. (2) Al<sub>2</sub>O<sub>3</sub> and Fe<sub>2</sub>O<sub>3</sub> are the mineral constituents occurring in next largest amounts, the former exceeding the latter by a considerable margin in most soils. (3) Either CaO or K<sub>2</sub>O is the fourth substance in order of quantity in the majority of soils. (4) There is generally less MgO than CaO, and less Na<sub>2</sub>O than K<sub>2</sub>O. (5) The percentages of Al<sub>2</sub>O<sub>3</sub>, Fe<sub>2</sub>O<sub>3</sub>, CaO, MgO, K<sub>2</sub>O, and Na<sub>2</sub>O are much more variable than is the percentage of SiO<sub>2</sub>. (6) The CaO and MgO content of soils from arid and semi-arid regions is appreciably higher than that of soils from humid regions. (7) The content of organic matter differs greatly. (8) P<sub>2</sub>O<sub>5</sub> and the other constituents not previously mentioned usually occur in fractions of a per cent.

# Composition of Soil Types

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TABLE 2  
CHEMICAL ANALYSES OF SEVERAL SOIL TYPES AND OF THE COLLOIDS EXTRACTED FROM THEM \*

| Soil Type and Location                      | Depth, inches | Constituent, per cent |                                |                                |      |      |                  |                   |                               |                | SiO <sub>2</sub><br>Al <sub>2</sub> O <sub>3</sub> + Fe <sub>2</sub> O <sub>3</sub> |
|---|---------------|-----------------------|--------------------------------|--------------------------------|------|------|------------------|-------------------|-------------------------------|----------------|---|
|   |               | SiO <sub>2</sub>      | Al <sub>2</sub> O <sub>3</sub> | Fe <sub>2</sub> O <sub>3</sub> | CaO  | MgO  | K <sub>2</sub> O | Na <sub>2</sub> O | P <sub>2</sub> O <sub>5</sub> | Organic matter |   |
| 1. Norfolk fine sandy loam, North Carolina  | 0-12          | 94.26                 | 2.08                           | 0.69                           | 0.07 | 0.03 | 0.03             | 0.01              | 0.01                          | 0.89           | 5.1   |
| 2. Bladen loam, North Carolina              | 0-8           | 89.96                 | 3.80                           | 0.72                           | 0.07 | 0.02 | 0.03             | 0.01              | 0.06                          | 2.74           | 4.4   |
| 3. Pecos sandy loam, New Mexico             | 0-9           | 66.39                 | 8.26                           | 2.48                           | 7.77 | 2.29 | 1.97             | 0.65              | 0.09                          | 1.25           | 8.2   |
| 4. Spearfish silt loam, South Dakota        | 0-12          | 68.71                 | 10.17                          | 3.51                           | 3.42 | 2.02 | 2.40             | 0.44              | 0.13                          | 3.26           | 7.7   |
| 5. Pullman silty clay loam, Texas           | 0-5           | 75.62                 | 10.56                          | 3.16                           | 0.67 | 0.78 | 2.20             | 1.06              | 0.10                          | 3.13           | 6.5   |
| 6. Pimo clay adobe, Arizona                 | 0-9           | 55.67                 | 16.60                          | 5.35                           | 5.38 | 2.52 | 2.47             | 0.74              | 0.19                          | 1.13           | 8.2   |
| 7. Cecil clay loam, Georgia                 | 0-9           | 83.81                 | 7.70                           | 2.97                           | 0.28 | 0.15 | 0.79             | 0.43              | 0.06                          | 0.56           | ...<br>1.48   |
| 8. Emmet fine sandy loam, Michigan          | 1-4           | 86.18                 | 5.67                           | 1.79                           | 0.54 | 0.18 | 2.61             | 0.74              | 0.05                          | 1.12           | 5.8   |
| 9. Carrington loam, Iowa                    | 0-12          | 77.28                 | 8.93                           | 2.89                           | 0.84 | 0.56 | 1.35             | 1.15              | 0.14                          | 3.94           | ...   |
| 10. Ontario loam, New York                  | 0-12          | 74.92                 | 10.47                          | 3.48                           | 1.23 | 0.98 | 1.25             | 2.15              | 0.15                          | 2.87           | 1.00  |
| 11. Orangeburg fine sandy loam, Mississippi | 0-10          | 93.66                 | 2.57                           | 0.93                           | 0.16 | 0.06 | 0.45             | 0.15              | 0.03                          | 0.77           | 0.50  |

*Composition of Colloids Extracted from the Above Soils*

|   |      |       |       |       |      |      |      |      |      |       |      |
|---|------|-------|-------|-------|------|------|------|------|------|-------|------|
| 1. Norfolk fine sandy loam, North Carolina  | 0-12 | 35.97 | 32.14 | 10.70 | 0.35 | 0.58 | 0.52 | 0.18 | 0.10 | 4.65  | 1.57 |
| 2. Bladen loam, North Carolina              | 0-8  | 45.76 | 31.33 | 3.20  | 0.57 | 0.30 | 0.20 | 0.17 | 0.30 | 6.41  | 2.83 |
| 3. Pecos sandy loam, New Mexico             | 0-9  | 44.08 | 18.36 | 6.45  | 8.42 | 3.83 | 2.34 | 0.26 | 0.26 | 2.19  | 3.33 |
| 4. Spearfish silt loam, South Dakota        | 0-12 | 45.29 | 22.91 | 6.40  | 3.34 | 4.08 | 2.33 | 0.25 | 0.25 | 5.84  | 2.85 |
| 5. Pullman silty clay loam, Texas           | 0-5  | 50.51 | 22.04 | 8.80  | 1.48 | 2.08 | 2.68 | 0.06 | 0.20 | 4.02  | 3.10 |
| 6. Pimo clay adobe, Arizona                 | 0-9  | 51.79 | 21.42 | 7.64  | 3.06 | 4.05 | 2.52 | T    | 0.21 | 1.45  | 3.34 |
| 7. Cecil clay loam, Georgia                 | 0-9  | 33.95 | 36.06 | 11.02 | 0.31 | 0.40 | 0.56 | 0.44 | 0.25 | 2.25  | 1.34 |
| 8. Emmet fine sandy loam, Michigan          | 1-4  | 41.10 | 19.67 | 6.46  | 1.54 | 1.92 | 2.09 | 0.26 | 0.29 | 15.18 | 2.94 |
| 9. Carrington loam, Iowa                    | 0-12 | 44.80 | 22.59 | 7.75  | 1.48 | 1.44 | 1.36 | 0.22 | 0.28 | 11.59 | 2.75 |
| 10. Ontario loam, New York                  | 0-12 | 38.01 | 23.30 | 10.84 | 1.37 | 2.34 | 2.23 | 0.14 | 0.44 | 17.13 | 2.13 |
| 11. Orangeburg fine sandy loam, Mississippi | 0-10 | 40.35 | 31.04 | 10.11 | 0.51 | 0.72 | 0.81 | 0.24 | 0.42 | 4.26  | 1.83 |

\* Compiled from U.S.D.A. *Bull.* 551, 1311; *Tech. Bull.* 170, 229, 502, 594.

The great variation in the composition of soils is illustrated by the data in Table 3. In obtaining these results Robinson and his associates analyzed forty-five soil samples from twenty-two locations in eight states.

TABLE 3

## EXTREMES IN COMPOSITION FOUND IN FORTY-FIVE SOILS \*

Constituent, range in per cent

| SiO <sub>2</sub> | Al <sub>2</sub> O <sub>3</sub> | Fe <sub>2</sub> O <sub>3</sub> | K <sub>2</sub> O | Na <sub>2</sub> O | CaO  | MgO  | P <sub>2</sub> O <sub>5</sub> | SO <sub>3</sub> |
|------------------|--------------------------------|--------------------------------|------------------|-------------------|------|------|-------------------------------|-----------------|
| 97.01            | 27.58                          | 16.23                          | 4.07             | 2.06              | 1.73 | 1.93 | 0.22                          | 0.34            |
| 44.15            | 1.19                           | 0.33                           | 0.02             | 0.01              | 0.01 | 0.01 | 0.03                          | 0.02            |

\* U.S.D.A. Bull. 551.

**Composition of Soil Separates and Classes.** As might be expected, the coarser soil separates contain a much higher proportion of minerals resistant to weathering, such as quartz, than do the finer separates. The content of several constituents such as P<sub>2</sub>O<sub>5</sub>, K<sub>2</sub>O, CaO, and MgO is appreciably higher in the finer than in the coarse separates, as the data in Table 4 show.

Because soil separates vary in composition and soil classes are made up of different proportions of the separates, it is logical to expect a difference in the chemical composition of the classes. Such difference is found unless some factor as age, parent material, climate, or management interferes. Usually sandy soils are higher in silica and lower in the other constituents, especially nitrogen, phosphorus, calcium, magnesium, and potash. The comparative nitrogen and phosphorus contents of a number of soil classes from New York illustrate this point (Table 5), although the variations in phosphorus content are not so consistent as are those for nitrogen.

A study of the composition of soil classes justifies the assumption that, in general, fine-textured soils are richer in total nutrients than are coarse-textured soils.

**Comparative Composition of Surface and Subsoils.** As soils are developed through the breaking down (weathering) of parent materials and through the building up of new materials, it is logical to expect an appreciable difference in both the chemical and mineralogical composition of soils and their parent materials. Usually the difference is greater between the *A* horizon, which has undergone the greatest amount of weathering, and the *C* horizon than between the *B* and *C* horizons. Also, more organic matter generally accumulates in the *A* horizon than in the *B* or *C* horizons. In some soils, however, the *B* horizon may exhibit a greater deviation in composition from the *C* than does the *A*.

TABLE 4  
COMPARATIVE NUTRIENT CONTENT OF SOIL SEPARATES \*

| Soil Origin and Number of Soils Analyzed                   | P <sub>2</sub> O <sub>5</sub> |      |      | CaO  |      |      | MgO  |      |      | K <sub>2</sub> O |      |      |
|--|-------------------------------|------|------|------|------|------|------|------|------|------------------|------|------|
|  | Sand                          | Silt | Clay | Sand | Silt | Clay | Sand | Silt | Clay | Sand             | Silt | Clay |
| Coastal Plains soils (7)                                   | 0.03                          | 0.10 | 0.34 | 0.07 | 0.19 | 0.55 | 0.09 | 0.14 | 0.61 | 0.37             | 1.34 | 1.76 |
| Soils developed from crystalline and metamorphic rocks (3) | 0.07                          | 0.22 | 0.67 | 0.50 | 0.82 | 0.94 | 0.48 | 0.86 | 1.24 | 1.60             | 2.37 | 2.86 |
| Glacial soils (10)   | 0.15                          | 0.23 | 0.86 | 1.24 | 1.30 | 2.69 | 0.54 | 0.88 | 1.80 | 1.72             | 2.35 | 3.08 |
| Soils developed from limestones and shales (5)             | 0.19                          | 0.17 | 0.49 | 7.55 | 6.82 | 6.67 | 0.44 | 0.52 | 1.84 | 1.46             | 1.95 | 2.67 |
| Arid soils (2)   | 0.19                          | 0.25 | 0.45 | 9.09 | 9.22 | 8.04 | 1.49 | 2.98 | 5.34 | 3.05             | 4.16 | 5.07 |
| General average  | 0.12                          | 0.18 | 0.61 | 2.61 | 2.57 | 3.07 | 0.47 | 0.77 | 1.70 | 1.41             | 2.15 | 2.78 |

\* Diameter of separates: sand, 2.00–0.05 mm.; silt, 0.05–0.005 mm.; clay, less than 0.005 mm. Compiled from data in "The Mineral Composition of Soil Particles," by G. H. Failyer, J. G. Smith, and H. R. Wade, U.S.D.A. Bur. Soils Bull. 54, 1908.

TABLE 5

## NITROGEN AND PHOSPHORUS CONTENTS OF SEVERAL NEW YORK SOIL CLASSES

| Soil Class          | Clay  | Silty |       |       | Loamy<br>and<br>Sands |
|---------------------|-------|-------|-------|-------|-----------------------|
|                     |       | Clay  | Loam  | Silt  |                       |
| Number of samples   | 21    | 31    | 58    | 47    | 19<br>3               |
| N content, per cent | 0.24  | 0.21  | 0.22  | 0.18  | 0.16<br>0.02          |
| Number of samples   | 21    | 31    | 61    | 48    | 19<br>3               |
| P content, per cent | 0.069 | 0.059 | 0.063 | 0.056 | 0.052<br>0.012        |

horizon because of concentration of certain constituents in it. Also, in some Podzols there is more organic matter in the *B* than in the *A* horizon because of the movement downward in colloidal state of this material. In general, in soils other than those of the tropics the *A* horizon is marked by an increase in organic matter content, a decrease in basic elements, especially Ca and Mg, and an increase in silica resulting from the loss of other constituents. The composition of the *A* and *C* horizons of a few soils is shown in Table 6.

**Soil Composition and Fertility.** Plants take all their mineral nutrients from the soil, and many obtain their nitrogen from the soil also. These minerals must come from the rocks and minerals in the soil, and they are made available for plant use through the processes of weathering. Accordingly, the percentages of the different mineral nutrients in a soil must depend on the quantities of the minerals which contain them. Furthermore, the degree of weathering which has taken place is important because some nutrients leach out rapidly as their parent minerals weather and when there is sufficient moisture to cause leaching. Also, because some minerals decompose very slowly in the soil, the chemical elements they contain become available for plant use in small amounts. On the other hand, some minerals may occur in small quantities in most soils, but because of their rapid rate of decay they set free their nutrient content quite readily. Plants also appear to require their nutrients in certain chemical combinations or forms before they can absorb them readily. As can be seen, the factors which determine the quantities of nutrients a given soil will supply to plants are quite involved and evidently the total quantities of nutrients a soil contains may not be a direct indication of the quantities a given crop can obtain from it. Various chemical methods have been devised in an effort to determine how much of the different nutrients plants can obtain from a given soil. Although progress has been made along this line, none of the methods is entirely successful, as is pointed out in Chapter 13.

TABLE 6

COMPOSITION OF SEVERAL SURFACE SOILS IN COMPARISON TO THAT OF  
THEIR PARENT MATERIALS

| Soil Number * | Horizon        | Constituent, per cent |                                |                                |       |      |                  |                               |        |
|---------------|----------------|-----------------------|--------------------------------|--------------------------------|-------|------|------------------|-------------------------------|--------|
|               |                | SiO <sub>2</sub>      | Al <sub>2</sub> O <sub>3</sub> | Fe <sub>2</sub> O <sub>3</sub> | CaO   | MgO  | K <sub>2</sub> O | P <sub>2</sub> O <sub>5</sub> | O.M. † |
| 1             | A              | 93.0                  | 2.9                            | 0.8                            | 0.13  | 0.05 | 0.05             | 0.04                          | 0.88   |
|               | C              | 75.5                  | 11.4                           | 4.4                            | 0.03  | 0.04 | 0.22             | 0.06                          | 0.17   |
| 2             | A              | 79.3                  | 10.9                           | 1.0                            | 0.33  | 0.06 | 5.15             | 0.26                          | 0.86   |
|               | C              | 70.5                  | 17.5                           | 1.0                            | 0.24  | 0.18 | 6.22             | 0.32                          | 0.19   |
| 3             | A <sub>1</sub> | 85.3                  | 7.6                            | 2.1                            | 1.46  | 0.72 | 2.05             | 0.15                          | 4.64   |
|               | C              | 66.1                  | 10.6                           | 3.1                            | 10.81 | 1.76 | 4.05             | 0.19                          | 0.68   |
| 4             | A <sub>0</sub> | 70.7                  | 9.6                            | 3.4                            | 1.66  | 0.96 | 2.11             | 0.26                          | 7.30   |
|               | C              | 68.6                  | 9.2                            | 2.8                            | 5.13  | 2.77 | 1.62             | 0.11                          | 0.00   |
| 5             | A <sub>1</sub> | 83.9                  | 6.5                            | 4.2                            | 0.64  | 0.49 | 3.47             | 0.24                          | 3.58   |
|               | C              | 79.6                  | 5.1                            | 3.4                            | 7.64  | 1.97 | 1.71             | 0.12                          | 0.38   |

\* Soil types: (1) Rushton fine sandy loam, North Carolina; (2) Durham sandy loam, Georgia; (3) Miami silt loam, Michigan; (4) Barnes silt loam, South Dakota; (5) Fox loam, Michigan.

† O.M. stands for organic matter.

In general, it can be said that soils which contain more than the average quantity of organic matter are more productive than are soils low in this constituent. Likewise, soils containing relatively high contents of calcium, especially in a readily soluble form, are usually productive. A high content of calcium carbonate, however, may have the reverse effect. And, as previously pointed out, soils containing enough silt and clay to place them in the loam, silt loam, and clay loam classes are usually more fertile than are the more sandy soils.

## THE COLLOIDAL FRACTION OF SOILS

It has been pointed out (p. 49) that the finer, more reactive part of the clay in soils is called colloidal clay. In most soils this fraction is made up of several materials, among the most common of which are (a) clay minerals, (b) organic matter, and (c) oxides of iron, aluminum, and silicon in various stages of hydration. The proportions of these different constituents vary considerably in different soils, as does also the nature of the clay minerals. For example, some quite sandy

soils may contain little colloidal material except organic matter, whereas in some clay soils the organic colloids are a small part of the total colloidal fraction. The quantity of colloidal matter in soils and some of its chemical properties are of much significance in the production of crops. Attention is called to these points in the following questions.

### Questions

1. How is the quantity of colloidal matter in soils determined?
2. How much colloidal material is there in soils?
3. What are clay minerals, and how do they differ in different soils?
4. What are the chemical properties of clay minerals?
5. What is the process of ion-exchange?
6. Is cation-exchange an important property in soils?
7. Are anions adsorbed by colloids?
8. How does ionic exchange function in the supplying of nutrients to plants?

**Determination of the Colloid Content of Soils.** The colloidal content of soils may be determined by various methods. As pointed out in Chapter 3, the hydrometer method may be used by allowing the dispersed soil suspension to stand for 24 hours and then determining the quantity of colloid in suspension by means of a specially designed hydrometer. Likewise, the pipette method is frequently used by drawing off a given volume of the suspension, after all but the colloidal material has settled, and evaporating it to dryness.

An older and less accurate procedure is based on the capacity of colloidal material to adsorb water and on the assumption that the colloidal fraction is the only portion of a soil that adsorbs water. For example, the quantity of moisture adsorbed from a saturated atmosphere over 3.3 per cent  $H_2SO_4$ , by a given weight of colloid extracted from a soil and spread in a thin layer, is determined. A similar determination is made on a sample of the soil itself. The colloidal content is then calculated by the equation

$$\frac{\% \text{ H}_2\text{O adsorbed by 1 gram of soil}}{\% \text{ H}_2\text{O adsorbed by 1 gram of colloid}} \times 100 = \% \text{ colloid content}$$

Another method is to determine the heat evolved when a given weight of oven-dried colloid is moistened with water. The heat-of-wetting of the soil is then determined. By substituting the heat-of-wetting in calories per gram of colloid and of soil in the above equation, the colloid content may be determined.

**Quantity and Composition of Colloids in Soils.** The colloidal content, like other soil constituents, varies greatly. Some sandy soils low in organic matter content contain a very small percentage of colloids, whereas some clay soils are very high in this constituent. In Table 7 is shown the colloidal content of several soils.

TABLE 7

## THE COLLOID CONTENT OF THE SURFACE HORIZON OF SEVERAL SOILS

| Soil and Location                          | Colloid Content, per cent | Soil and Location               | Colloid Content, per cent |
|--|---------------------------|---------------------------------|---------------------------|
| Miami silt loam, Indiana                   | 15.8 *                    | Pierre clay, South Dakota       | 57.4 †                    |
| Chester sandy loam, Maryland               | 14.2 *                    | Ruston fine sandy loam, Georgia | 7.3 †                     |
| Cecil sandy loam, Georgia                  | 10.2 *                    | Bladen loam, North Carolina     | 11.4 †                    |
| Amarillo silty clay loam, Texas            | 23.9 †                    | Hillsdale sandy loam, Michigan  | 15.7 ‡                    |
| Nipe clay, Cuba                            | 50.2 †                    | Yolo loam, California           | 48.5 ‡                    |
| Davidson clay loam, North Carolina         | 25.9 †                    | Marion silt loam, Minnesota     | 45.4 ‡                    |
| Orangeburg fine sandy loam, North Carolina | 3.0 †                     | Brookston clay loam, Michigan   | 33.4 ‡                    |
| Pima clay, Arizona                         | 59.2 †                    | Sandy loam, Rhode Island        | 17.7 ‡                    |

\* Determined by the water-adsorption method.

† Determined by the pipette method.

‡ Determined by the heat-of-wetting method.

The composition of colloidal material does not follow that of the soils containing it as is seen by the data in Table 2. It will be noted that the  $\text{SiO}_2$  content of the colloidal fraction is much lower than that of the corresponding soil, whereas the  $\text{Fe}_2\text{O}_3$  and  $\text{Al}_2\text{O}_3$  content is much higher. Also, the colloids contain considerably more  $\text{CaO}$  and  $\text{MgO}$  than do their parent soils in most soils. The higher percentage of  $\text{P}_2\text{O}_5$  and organic matter in the colloids than in the soils is also noteworthy. The very high organic content in some of the colloids from soils which contain a medium to low percentage of this material is of interest.

**Clay Minerals.** As pointed out on pages 32 and 50, the clay minerals are compounds developed in the soil. They are not fragments of minerals which were in the parent material. These minerals have definite crystal structures by which they may be identified through use of the X-ray and electron microscope. The climatic conditions under which weathering occurs have much to do with the relative proportions of the different clay minerals developed from the products of weathering. For example, under warm, humid conditions a greater proportion of minerals of the kaolinite group than of the montmorillonite

minerals is developed. In cool, moist climates the condition is reversed. As each of the groups of minerals has specific chemical and physical properties, the proportion of each mineral present influences the characteristics and reactions of the soils containing them.

These minerals are made up of plates of  $\text{Al}_2\text{O}_3$  and  $\text{SiO}_2$  and are generally plate-like in shape, and so they expose a very large amount of surface per unit of weight. For example, Baver,<sup>1</sup> quoting Ostwald, points out that a spherical particle with a volume of 1 cc. exposes 4.836 sq. cm. of surface, a cubical particle of equal volume has a surface of 6.0 sq. cm., and a plate-shaped particle with a height of 1 mm. exposes 21.121 sq. cm. of surface. Naturally, as the height of the plate decreases, the surface area increases very rapidly. This large amount of surface has much to do with the chemical and physical effects of clay minerals in soils. Furthermore, the capacity of members of the montmorillonite group of minerals to expand on wetting greatly influences their chemical and physical activities. In order to emphasize the difference in make-up of the principal clay mineral groups, the composition of some of them is repeated here. The montmorillonite group,  $(\text{OH})_4\text{Al}_4\text{Si}_8\text{O}_{20} \cdot \text{XH}_2\text{O}$ , is characterized by a high  $\text{SiO}_2$  content, as it contains two  $\text{SiO}_2$  plates, and by a low content of OH groups. On the other hand, the kaolinite group,  $(\text{OH})_8\text{Al}_4\text{Si}_4\text{O}_{10}$ , contains one plate of  $\text{SiO}_2$  and hence is low in silica but has a much larger number of OH groups. In the illite group the replacement of Al by Fe and sometimes Mg leads to considerable variation in composition,  $(\text{OH})_4\text{K}_x(\text{Al}_4 \cdot \text{Fe}_4 \cdot \text{Mg}_4\text{Mg}_6)(\text{Si}_{8-x} \cdot \text{Al}_x)\text{O}_{20}$ . Note that the content of OH is low and that of  $\text{SiO}_2$  is high as in montmorillonite. In general, the greater the ratio of silica ( $\text{SiO}_2$ ) to sesquioxides ( $\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3$ ), the greater is the chemical activity of the colloid mineral. This ratio is calculated as follows:

$$\frac{\% \text{ SiO}_2}{\text{Molecular wt. of } \text{SiO}_2 (60.1)} = A$$

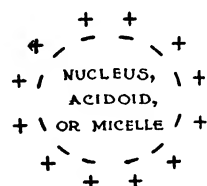
$$\frac{\% \text{ Al}_2\text{O}_3}{\text{Molecular wt. of } \text{Al}_2\text{O}_3 (101.9)} = B$$

$$\frac{\% \text{ Fe}_2\text{O}_3}{\text{Molecular wt. of } \text{Fe}_2\text{O}_3 (159.7)} = C$$

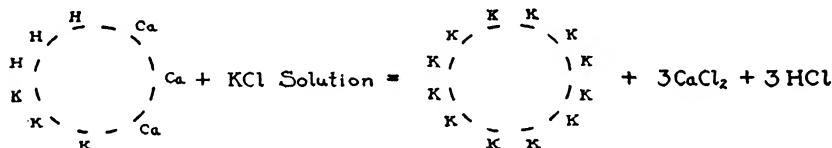
$$\frac{A}{B + C} = \text{silica:sesquioxide ratio}$$

<sup>1</sup> L. D. Baver, *Soil Physics*, 2nd edition, John Wiley & Sons, Inc., 1948.

**Chemical Properties of Clay Minerals.** Chemically many clay minerals are similar to the negative radical of an acid, as  $\text{PO}_4^{4-}$  or  $\text{SO}_4^{2-}$ . In other words, they are complex, negatively charged nuclei which will move toward the positive electrode when submitted to the action of an electric current. They may be pictured roughly as shown in the diagram. The inner layer of negative charges is part of the wall of the particle. The outer layer is at a distance of molecular dimensions from the inner layer. In the diagram the inner layer is of negative charges and the outer positive. In some colloids the situation is reversed although the arrangement shown is the usual one. The outer layer may be replaced by other ions of a similar charge as  $\text{K}^+$ ,  $\text{Na}^+$ ,  $\text{Ca}^{++}$ , and  $\text{Mg}^{++}$ . As might be expected, the behavior of a clay particle is greatly influenced by the nature of the ions in the outer layer.



**Ion-Exchange.** The fixation or adsorption of an ion by the colloidal nucleus, acidoid or micelle and the accompanying release of one or more ions held by the nucleus is termed *ion-exchange*. For example, assume that the nucleus has one-half of its capacity satisfied with Ca ions, one-quarter with K ions, and one-quarter with H ions. Diagrammatically the situation would be as shown in the diagram below. Now suppose



that the colloidal material is treated with a strong solution of KCl. First the K ions replace the Ca ions on the complex, and they in turn combine with the Cl ions to form  $\text{CaCl}_2$ . Upon continued treatment with the KCl solution the H is replaced. Thus by suitable procedure all the exchangeable bases on the nucleus may be replaced by any desired ion.

The efficiency with which ions will replace each other is determined by such factors as (a) relative concentration or numbers of the ions, (b) the number of charges on the ions, and (c) the speed of movement or activity of the different ions. The first factor is an application of the well-known chemical law of mass action. The greater the number of charges carried by the ion the greater is its efficiency, other factors being the same. The speed or activity of an ion is primarily a function of its size, in which the degree of hydration must also be considered.

If we consider the ions Li, Na, K, and Rb, we find that they are listed in the ascending order of size, and hence we would expect their efficiency of replacement to be Li Na K Rb. In reality, however, the Li ion associates itself with so many water molecules that its speed is much reduced, and, furthermore, it cannot get near the micelle because of the shell of water. Likewise, the Na ion is more highly hydrated than the K ion. The result is that the order of replacement is reversed: Rb K Na Li. A listing of the more common ions in descending order of replacing power is: H Sr Ba Ca Mg Rb K NH<sub>4</sub> Na Li.<sup>2</sup>

It has been noted that organic material may make up an appreciable portion of the colloidal content of soils; in fact, it may constitute the major part of the colloids in sandy soils. The ion-exchange capacity of organic colloids therefore becomes significant. It is noteworthy that, although the exchange capacity of organic colloidal materials varies greatly, as does that of clay minerals, on the whole it runs from five to seven times that of mineral colloids on a weight basis.

**Cation-Exchange of Soils.** The taking up and giving off of positively charged ions by a soil or other substance is referred to as cation- or base-exchange. The total cation-exchange capacity of a soil is expressed by the milliequivalents (m.e.) of ions 100 grams of soil will adsorb. When the complex has its entire capacity satisfied by H<sup>+</sup>, it is said to be unsaturated. When other ions satisfy a part of the exchange capacity, the degree of saturation is expressed as the percentage of the total capacity satisfied by such ions. When all the capacity is satisfied with basic ions, it is called completely saturated or base-saturated.

If a soil has a large cation-exchange capacity and this capacity is entirely satisfied with basic ions such as Ca, Mg, K, and Na, a large quantity of H ion is required to change appreciably the chemical and physical characteristics of the colloid, and the soil is said to be highly buffered. Likewise, if the colloid were largely saturated with H ion, it would take much Ca, Mg, and the like to change its characteristics. On the other

<sup>2</sup> There are certain difficulties in accepting the idea that the hydration of ions is a determining factor in cation-exchange; yet there is much evidence in support of this theory, and it is believed to be a useful concept. However, according to Grim, cation exchangeability cannot be explained satisfactorily on the basis of cation hydration. He points out that some cations, previously thought to be highly hydrated, probably do not hydrate at all and that other cations hydrate to a lesser degree than has been assumed. [R. E. Grim, "Modern Concepts of Clay Minerals," *Journal of Geology*, Vol. 50, 1942, pp. 225-275.]

TABLE 8

## IONIC RADII, HYDRATION, VELOCITY, AND EXCHANGE EFFICIENCY OF SEVERAL IONS

| Ion             | Radii of Ions,<br>Angstroms<br>( $10^{-8}$ cm.) |          | Velocity of<br>Migration,<br>microns per sec.<br>per volt per cm. | No. of<br>Charges<br>Carried | Order of Cation-<br>Exchange<br>Efficiency |
|-----------------|---|----------|---|------------------------------|--|
|                 | Dehydrated                                      | Hydrated |   |                              |  |
| Li              | 0.78  | 10.03    | 3.45  | 1                            | 8th  |
| Na              | 0.98  | 7.90     | 3.31  | 1                            | 7th  |
| K               | 1.33  | 5.32     | ....  | 1                            | 5th  |
| NH <sub>4</sub> | 1.43  | 5.37     | 3.48  | 1                            | 6th  |
| Rb              | 1.49  | 5.09     | 3.25  | 1                            | 4th  |
| H               | ....  | ....     | 2.84  | 1                            | 1st  |
| Ca              | 1.06  | ....     | 3.27  | 2                            | 2d   |
| Mg              | 0.78  | ....     | 3.18  | 2                            | 3d   |

hand, a soil having a small exchange capacity will have the characteristics of the colloidal fraction modified by the adsorption of a relatively small amount of base or of H ion. Such a soil is said to be lightly buffered.

The expanding 2:1 crystal lattice of the montmorillonite minerals gives them a much greater cation-exchange capacity than have the kaolinite minerals.

**Anion-Exchange.** Although cation-exchange in soils has been studied extensively and its importance in supplying nutrients to plants is recognized, the phenomenon of the exchange of anions in the colloidal complex has received less attention. It has been established that such exchange does occur, though apparently to a far less extent than in cations. Anions may replace the OH groups in the clay minerals, and, as these groups are much more plentiful in the kaolinite minerals than in the montmorillonite minerals, the kaolinite minerals are considered the seat of most anion-exchange. It is possible, however, that considerable anion-exchange may take place in the amorphous colloidal material in the soil. This phenomenon needs much more investigation to determine its relative significance in soil reactions and plant nutrition.

**Soil Colloids as Sources of Plant Nutrients.** It has been noted that the H ion is the most active in cation replacement. Hence, a de-

velopment of H ions in the soil would lead to the replacement of other cations on the colloidal complex, and these replaced ions would then be available for use by plants. Fortunately, there is a constant liberation of H ions in the soil during the growing season. This comes about through the production of  $\text{CO}_2$  from the decay of organic matter and the excretion of the gas from living cells by the process of respiration.

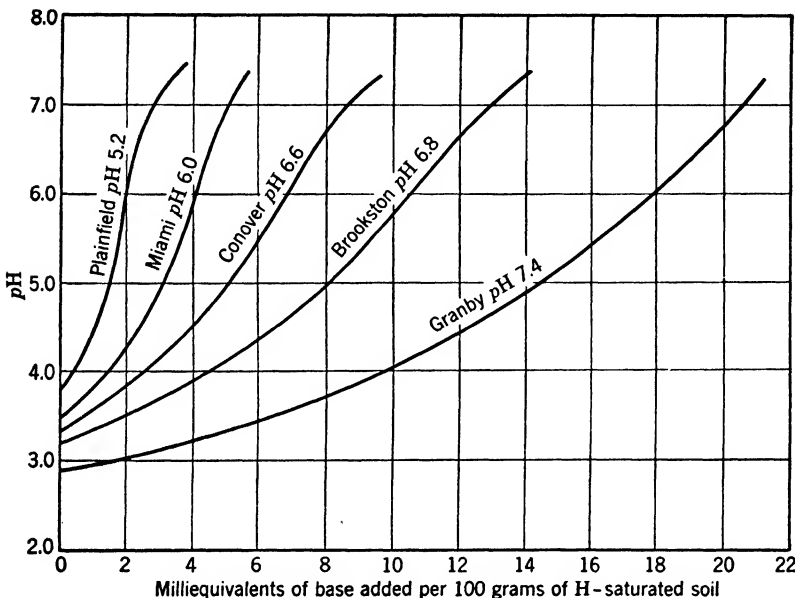


FIG. 28. Electrometric titration curves showing the different amounts of base needed to increase the  $\text{pH}$  of soils having different degrees of buffering. All soils were H-saturated when titration was started.

The  $\text{CO}_2$  combines with water, forming  $\text{HCO}_3^-$  ion and H ion, which may then replace other cations for plant use.

It is also possible for anions such as  $\text{PO}_4^{3-}$  to be liberated from colloidal material and then be taken up by plants. Some workers think that adsorbed  $\text{PO}_4^{3-}$  is an important source of this nutrient for plant growth. On the other hand, some anions, as  $\text{NO}_3^-$ , are adsorbed to a small extent if at all by soil colloids.

When commercial fertilizers, manures, and lime are applied to soil, it is possible for the colloids to take up considerable quantities of various ions to be liberated later for use by crops. In general, soils with a relatively high ion-exchange capacity are fertile and have good lasting quality for crop production; that is, they are depleted more slowly than are soils

with low ion-exchange capacity. The colloidal complex may therefore be considered a storehouse for plant nutrients.

#### OTHER CHEMICAL PROPERTIES OF SOILS

A soil might be considered a huge laboratory in which innumerable physical, chemical, and biological processes are taking place. Few, if any, of these processes are independent, but each is influenced by many others. A few of the more important physical and chemical processes have been discussed, and attention is called to the biological phase of soil science in Chapter 9. Only a limited number of chemical reactions and processes are mentioned in this chapter, and it should be recalled that several more have been discussed in the sections dealing with weathering and soil formation. Likewise, many chemical properties of soils center around soil reaction (acidity and alkalinity) and these are taken up in the following chapter. Furthermore, the addition of fertilizer and lime and the decay of organic matter result in numerous reactions concerned with the fixation, liberation, and utilization of nutrients, and these are considered in their proper places. The discussion of chemical properties and processes of soils presented in this chapter is, accordingly, very limited although highly significant.

# 6

## Soil Reaction

Soils may be acid, neutral, or alkaline in reaction. The reaction of soils is significant in crop-production and soil-management practices because the various degrees of soil reaction are produced by the chemical conditions which exist in soils, each set of chemical conditions causing its corresponding degree of soil reaction. Each degree of soil reaction or set of chemical conditions in soils affects plant growth in a certain way owing to either a depressed solubility of some elements or to an increased solubility of others. The chemical conditions which accompany the different degrees of soil reaction, therefore, may be favorable to the growth of some crops, unfavorable to others, and in still other cases they may affect plant growth very little. Soil reaction, then, may be considered a *symptom of the particular chemical conditions which caused it*, and hence it may be used to indicate the possible effect of these conditions on plant growth. It is useful in diagnosing the fertility of soils. A knowledge of the conditions which cause different soil reactions, therefore, is of value to the student of soil science.

A Virginia farmer named Edmund Ruffin (1794–1865) was probably the first man in the United States to sense the prevalence of acidity and the need for lime in soils of eastern United States. In the latter part of the nineteenth and early years of the present century Dr. Wheeler, of the Rhode Island Experiment Station, made soil tests for acidity over many of the eastern and central states and by his writings and lectures gave much publicity to the widespread need for liming soil.

The study of soil reaction and of the effects on plant growth of the chemical conditions accompanying various soil reactions may be taken up under three general heads or objectives.

### Objectives

- A. Soil acidity and conditions which produce acid soils.
- B. Properties of acid soils which affect plant growth.

- C. Conditions of development and general effect of neutral and alkaline soils on the growth of plants.

### SOIL ACIDITY AND CONDITIONS GIVING RISE TO ACID SOILS

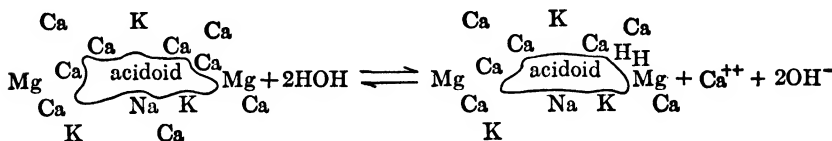
There is a natural tendency for soils to become increasingly acid in humid climates, and this trend is accelerated when soils are put under cultivation. Large areas of soil were acid when the land was cleared in preparation for farming, but the store of available plant-food elements in the virgin soils and the supply of bases liberated through rapid decay of the organic matter largely offset the detrimental effects of the acid condition. Consequently, farmers did not become conscious of the need of the soil for lime until clover began to fail. As most of the highly prized leguminous crops do not thrive on soils of strong acidity and as the yields of many other crops are decreased, the need of acid soils for lime becomes a very significant problem in the agriculture of humid regions. A series of questions commonly asked about soil acidity is listed.

#### Questions

1. What is meant by an acid soil?
2. Into what two types may soil acidity be classified?
3. What kind of rocks give rise to acid soils?
4. How are acid soils developed from non-acid minerals?
5. What farm practices tend to increase soil acidity?
6. Where are acid soils found?
7. How is the acidity of soils expressed?
8. How acid and how alkaline may soils be?
9. Does the reaction of soil vary during the year?
10. How may soils be made more acid?
11. What is meant by buffering in soils?
12. In what ways may soil acidity be determined?
13. Does a soil acidity test tell how much lime is needed?

**Acidity in Soils.** Soil acidity is primarily a function of the colloidal fraction of soils, but soluble acids may contribute to the condition to a greater or lesser extent. It has been estimated that at least 95 per cent of the acidity of mineral soils is due to the reaction of the colloidal material. As previously pointed out, much of the colloidal material of soil reacts as a complex acid radical of very low solubility, around which are clustered positive ions at varying distances from the particle surface. These cations dissociate to some extent, and their places on the acidoid are taken by H ions from the water, as shown in the illustration

below. The OH ions so formed produce an alkaline reaction in the soil solution. Although the H ions in the complex will also dissociate to some extent, there is reason to believe that they are held quite close to the acidoid, while other cations may be scattered at greater distances in the field of attraction.



When the numbers of H ions and OH ions in the solution around the nucleus are equal, a neutral reaction exists. When the number of H ions exceeds the number of OH ions, the reaction of the soil is acid.

Traces of inorganic acids, such as  $\text{HNO}_3$ ,  $\text{HNO}_2$ ,  $\text{H}_2\text{SO}_4$ , and  $\text{H}_3\text{PO}_4$ , may be present, and in unusual soils which contain  $\text{FeS}_2$  a considerable concentration of  $\text{H}_2\text{SO}_4$  may exist. Carbon dioxide is present in most soils and with  $\text{H}_2\text{O}$  produces a slightly acid reaction. Other organic acids, such as acetic, citric, and oxalic, are liberated during the decay of fresh plant tissue and may exist in the soil for a short period. Acids of this kind, however, are readily decomposed by soil microorganisms in well-aerated soils and so have only a temporary existence in the soil. Although inorganic and organic acids may contribute to some extent to the acidity of mineral soils, the reaction is due primarily to the relative proportion of H ions and other cations on the colloidal complex.

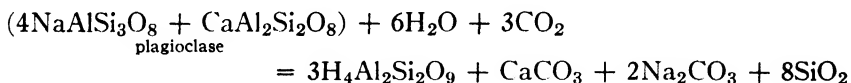
Studies have indicated that when the complex is saturated with divalent ions, such as  $\text{Ca}^{++}$  or  $\text{Mg}^{++}$ , a slightly alkaline reaction results and, furthermore, that a soil with reaction approaching neutrality contains very little exchangeable hydrogen.

**Types of Soil Acidity.** For a soil to give an acid reaction there must be an excess of H ions over OH ions in the solution surrounding the colloidal complex. The concentration of these H ions can be readily measured and constitutes the *active* acidity of the soil. The concentration of these dissociated H ions is in equilibrium with those adsorbed on the complex. Should any process, such as the addition of lime or leaching, remove or neutralize a portion of the active H ions, more would immediately dissociate from the complex to restore the equilibrium. The H ions on the acidoid, therefore, constitute a reserve supply which is known as the *reserve* or *potential* soil acidity.

**Soils from Acid Rocks.** Rocks which contain an excess of quartz or of silica compared to their content of basic minerals or of basic

elements are classed as acid rocks. Granite and rhyolite are examples. This classification is based on the fact that  $\text{SiO}_2$ , combined with various proportions of water, forms the different silicic acids of which orthosilicic acid,  $(\text{H}_2\text{O})_2\cdot\text{SiO}_3$ , and trisilicic acid,  $(\text{H}_2\text{O})_2\cdot 3\text{SiO}_2$ , are examples. When rocks which are deficient in bases are disintegrated or decomposed in the process of the accumulation of soil material and soil development, the resulting material is acid even though no loss of base has taken place during the process. There are considerable areas of sandy soils which have been produced from acid rocks.

**Acid Soils from Non-Acid Minerals.** Aside from the oxides and carbonates, most minerals are compounds of basic elements with some of the silicic acids or with aluminosilicic acid. As these compounds are broken down by the chemical processes of weathering, at least a part of the bases passes into the carbonate or bicarbonate forms, which have an appreciable solubility and are leached away when rainfall is sufficient. On the other hand, the negative or acid radicals of the compounds pass over into hydrated aluminum silicate and acid salts of silicic acid usually with the elimination of some free  $\text{SiO}_2$ . The following reaction is typical:



The acid salts of silicic acids give acid reactions just as monocalcium phosphate,  $\text{Ca}(\text{H}_2\text{PO}_4)_2$ , and potassium hydrogen sulfate,  $\text{KHSO}_4$ , react acid. The hydrated aluminum silicates are really aluminosilicic acids which are relatively insoluble and so accumulate in the soil. They are chiefly crystalline in structure and constitute some of the so-called clay minerals. The result of the weathering process is the accumulation of material which grows constantly more deficient in bases.

During the decomposition of the soil minerals are developed the clay minerals, some of which may have a high ion-exchange capacity. As bases are made soluble through mineral decomposition, they are adsorbed by the organic colloids and clay minerals, giving a complex with a high degree of saturation and an alkaline reaction. As mentioned in the preceding chapter, considerable  $\text{CO}_2$  is liberated in the soil through the decay of organic matter and by the respiration of plant roots. Carbon dioxide combines with water to form the bicarbonate and hydrogen ions.  $\text{CO}_2 + \text{HOH} = \text{HCO}_3^- + \text{H}^+$ .

It will be remembered that the  $\text{H}$  ion is the most effective in cation replacement, and hence the bases on the colloidal acidoid are con-

stantly being dislodged by the hydrogen ions which are continuously generated in the soil during the growing season. The bases so replaced may be used by plants, or they may be leached out of the soil. It will be seen, then, that here is another force which continually operates to lower the base content of the soil and make it acid. Because of the small exchange capacity of sandy soils, they become acid before the heavier soils, other factors being equal.

**Farm Practices that Increase Acidity.** Under virgin conditions soils are covered with vegetation throughout the year, and during growth periods the use of water by the plants materially reduces the quantity which percolates through the soil and removes soluble bases. The intake of nutrients by plants during virtually all the warm part of the year also reduces losses by leaching. Furthermore, all the basic nutrients taken up by the vegetation are sooner or later returned to the soil. It is to be expected, then, that development of acidity through loss of bases is a relatively slow process in virgin soils.

When the land is used for farming, a rather different set of conditions prevails. Unless cover crops are used extensively, under most rotations the soil is devoid of vegetative cover during many months of growing weather. Even with a carefully planned use of cover crops, the soil is unprotected from leaching during the fitting of the seedbeds. Fall plowing leaves the soil in excellent condition for loss of bases by leaching during late fall and early spring. Summer fallowing also leads to much loss by leaching. Through the harvesting of crops, most of the nutrients used by the plants are removed and a comparatively small proportion of them is returned in manure. It appears, therefore, that farming hastens the development of acidity and that practices which leave the soil free of growing vegetation and so subject to leaching are particularly objectionable.

Certain fertilizers tend to increase soil acidity through the development of mineral acids. This is particularly true of sulfate of ammonia. Other fertilizers, as calcium cyanamide, nitrate of soda, and calcium nitrate, tend to correct acidity. The effect of fertilizers on acidity will be discussed more fully in Chapter 15 which deals with fertilizers.

**Distribution of Acid Soils.** Acid soils will develop most rapidly in areas where rainfall is sufficiently abundant to leach away rapidly the bases liberated by ionic exchange and mineral decomposition. As the amount of water percolating through the soil diminishes, the rate at which acidity develops decreases. There are some soils in which acidity has become so pronounced and widespread as to require the application

of considerable quantities of lime in order to permit the satisfactory growth of acid-sensitive legumes. These soils occur primarily east of a line drawn roughly north and south along the eastern boundary of Kansas. Many areas of soil west of this boundary are improved by liming, and in fact quite strongly acid soils are found near the mountains adjoining deserts. Large acreages of land east of the boundary line are adequately supplied with bases, particularly in parts of the tall-grass prairie, in shallow or young soils derived from limestone, and in young soils developing on poorly drained locations and recently exposed lake beds. There are also areas in which leaching has not yet been sufficient to remove the limestone from the glacial till or other parent material.

Not all soils giving a moderately low *pH* value are in need of lime. In some soils of moderate to high acidity the exchange complex yet contains an abundant supply of bases which may supply the demand of crops for these nutrients.

**Expression of Soil Acidity.** There are several systems of expressing the acidity of soils. Within the last few years it has become customary to base the expression of acidity on the concentration of the dissociated H ions, that is, on the active acidity. This tendency has given rise to use of the *pH* values so frequently seen. The *pH* scale is derived as follows: Water is a neutral substance with equal concentrations of H and OH ions. The ionization constant of water is  $10^{-14}$ . As concentrations of  $H^+$  and  $OH^-$  are equal, each must be  $10^{-7}$ .

$$\frac{\text{Conc. } H^+ \times \text{conc. } OH^-}{\text{Conc. } H_2O} = 10^{-14}$$

As concentration of  $H_2O$  (conc.  $H_2O$ ) may be regarded as a constant and almost equal to the molar concentration per liter of water it may be ignored, and we may write

$$\text{Conc. } H^+ \times \text{conc. } OH^- = 10^{-14}$$

Considering only the concentration of the H ion because it is the cause of the acid reaction, we have

$$\text{Conc. } H^+ = 10^{-7} = 0.0000001 \text{ gram H ions per liter}$$

The reciprocal of this is  $1/0.0000001 = 10,000,000$ , the log of which is 7. Pure water is therefore said to have a *pH* of 7, and as water is neutral any solution with a *pH* of 7 is said to be neutral. It is seen from

the above that the *pH* value is the logarithm of the reciprocal of the H-ion concentration.

If the concentration of H ions in a solution were 0.000001 gram per liter ( $10^{-6}$ ), the *pH* would be 6. As the concentration of H ions in this solution is 10 times that in water, it is seen that a solution with *pH* 6 is 10 times as acid as a solution with *pH* 7. It should never be forgotten that *pH* values are logarithms.

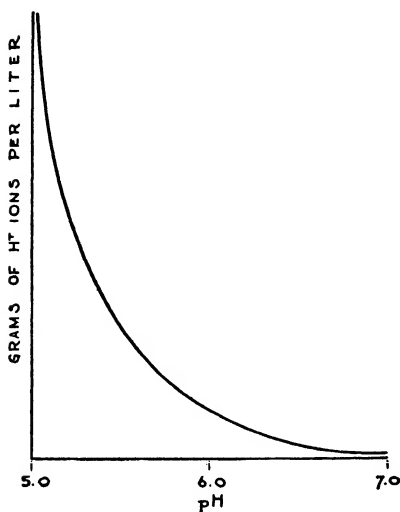


FIG. 29. In the *pH* scale the concentration of hydrogen ions is multiplied by 10 for each unit decrease in the scale. In the curve it will be noted how rapidly the hydrogen-ion concentration increases with *pH* decrease. For a *pH* value of 4 the line representing grams of hydrogen per liter would be over 4 feet long.

measures of total acidity. For example, two soils may have the same *pH* value but quite different total acidities or, in other words, may require decidedly different quantities of lime entirely to saturate the colloidal complex. Let us suppose that one soil, because of the quantity or nature of the colloidal material it contains, has a cation-exchange capacity of 100 m.e., while another soil has a capacity of only 50 m.e. If the capacity of each soil is one-half supplied with calcium and one-half with hydrogen, they would have about the same *pH*. Despite this fact, it would require twice as much calcium to saturate the complex of the first soil as it would to saturate the complex of the second soil. Be-

As the product of the concentration of the H and OH ions is always equal to  $10^{-14}$ , the concentration of the OH<sup>-</sup> is known if we express the *pH* value. Thus *pH* 7 indicates a concentration of both H and OH ions of  $10^{-7}$  ( $10^{-7} \times 10^{-7} = 10^{-14}$ ). A *pH* of 6 would then indicate a concentration of OH ions of  $10^{-8}$  ( $10^{-6} \times 10^{-8} = 10^{-14}$ ). On the other hand a *pH* of 8 would require a concentration of OH ions of  $10^{-6}$ , which is 10 times greater than  $10^{-7}$ , the concentration of OH ions in a neutral solution. This solution therefore is alkaline. It is seen then that *pH* values above 7 indicate alkaline solutions.

Although *pH* values express the active acidity or intensity of acidity in a soil, they do not indicate the reserve acidity and hence are not

cause of this situation the relative acidities of different soils are sometimes expressed in terms of the quantity of limestone ( $\text{CaCO}_3$ ) required to bring them to given  $p\text{H}$  values, as 6.5 or 7.0.

In other instances the degree of saturation is given in terms of percentage of the total cation-exchange capacity in order to express acidity or lime requirement. Thus a soil may be said to have an exchange capacity of 80 m.e. with 75 per cent saturation.

**Range of Reaction in Soils.** Unless there is an abnormal development of strong acid such as  $\text{H}_2\text{SO}_4$ , it is unusual to find mineral soils with a  $p\text{H}$  reading of less than 3.5; in fact,  $p\text{H}$  4.0 might more correctly be given as the lower limit frequently encountered. In humid regions it is unusual to find a soil with a  $p\text{H}$  of more than 7.5 or 8, but in arid regions where soluble salts of sodium ( $\text{Na}_2\text{CO}_3$ ) may accumulate, a  $p\text{H}$  of 9.5 to 11.0 is sometimes attained. The usual  $p\text{H}$  of agricultural soils in the humid region ranges approximately from 5.0 to 6.8.

In discussing soil acidity with those who are not familiar with the  $p\text{H}$  scale, it is customary to use descriptive terms to indicate the acidity and comparative lime requirement of soils. The scheme presented in Chart 3, with corresponding  $p\text{H}$  values, is representative.

**Fluctuations in Acidity.** It has frequently been observed that  $p\text{H}$  values of samples taken from a soil area at different times in the year show considerable variation. Several investigators have studied  $p\text{H}$  changes in soils by systematically sampling them during the growing season. Results show that acidity tends to increase from spring to mid-summer in humid regions. The increase is accentuated by periods of drought and may amount to an entire  $p\text{H}$  unit. Rainy periods decrease acidity, and, as winter approaches, the soil gradually returns to the  $p\text{H}$  observed in the early spring. An increase in  $p\text{H}$  has also been observed to accompany the thawing of frozen soil. The seasonal decrease in soil  $p\text{H}$  has been found to correlate roughly with an increase in soluble salt content, particularly of nitrates. A rapid growth of plants with accompanying absorption of nutrients restricts the accumulation of salts and so modifies the increase in acidity. The dehydration of colloidal material has also been suggested as a cause contributing to increased acidity in warm, dry weather. Fertilizers appear to have little influence on the seasonal fluctuation of soil  $p\text{H}$  unless they contain  $(\text{NH}_4)_2\text{SO}_4$  or other nitrogen compounds which produce acids in the soil.

The tendency of soils to increase in acidity in summer may be of small significance except in soil of low clay content, in which comparatively small applications of lime induce considerable changes in  $p\text{H}$ . When such soils are used for cropping systems in which small variations

### Chart 3. A Summary of Soil Reaction, Lime Requirement, and Associated Plant-Nutrient Relationships

| pH                   | 4.0                                       | 4.5           | 5.0  | 5.5   | 6.0     | 6.5                   | 6.7                                     | 7.0               | 8.0                                  | 9.0                                     | 10.0 | 11.0   |
|----------------------|---|---------------|--|---|---------|-----------------------|---|-------------------|--------------------------------------|---|------|--|
| Acidity              | Very strongly acid                        | Strongly acid | Moderately acid                            | Slightly acid                               | Neutral | Weakly alkaline       | Alkaline                                | Strongly alkaline |                                      |   |      |  |
| Lime requirements    | Lime needed for crops requiring acid soil |               |  | Lime needed for all but acid-tolerant crops |         |                       | Lime generally not required             |                   |                                      | No lime needed                          |      |  |
| Occurrence           | Rare                                      | Frequent      | Very common in plow soil in humid climates |   |         | Plow soils when limed |   |                   | Common in sub-humid and arid regions |   |      | Black alkali occurs in limited areas in arid regions |
| Fertility conditions | Phosphates fixed                          |               |  | Phosphates soluble                          |         |                       | Phosphate fixed                         |                   |                                      | Desirable bacterial activity            |      |  |
|                      | Lime and potash leach                     |               |  | Iron, aluminum, and manganese are soluble   |         |                       | Lime present in optimum quantities      |                   |                                      | Excess boron may occur in limited areas |      |  |
|                      | Fungi thrive                              |               |  | Bacteria languish                           |         |                       | Excess boron may occur in limited areas |                   |                                      | Nitrogen freely fixed                   |      |  |

in acidity may affect quality, yield, or disease resistance of the crops, seasonal variations in acidity are more noteworthy.

Air drying of surface soil samples appears to change pH values very little in acid soils but to reduce the alkalinity of alkaline soils. The pH of subsoil samples decreases appreciably with air drying.

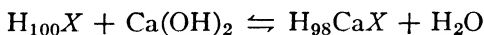
Because the mineralogical composition of a body of soil is not uniform, some variation in the pH of samples taken at various places is to be expected. Likewise the uneven distribution of applied manure and of manure dropped by animals may cause local variation in acidity in a field for several years. The burning of weeds, grass, or brush, the digging of a fence post hole, or the removal of stumps or large rocks may also cause a soil sample to be unrepresentative of a field. As a result, several samples from a field should be taken for testing. In fields with rolling topography there frequently is considerable difference in reaction of soil in the depressions and on the higher ground. The soil occupying the lower sites is usually of higher pH than that on the knolls and slopes.

**Ways of Increasing Acidity.** There are a number of plants which grow well only in soils that are strongly acid. Some of those most commonly grown are cranberry, blueberry, rhododendron, and azalea. At times it becomes desirable to increase the acidity of the soil for the growing of some of these plants. Several procedures used to accomplish this purpose are (1) the addition of ground sulfur, which oxidizes in the soil to sulfuric acid; (2) the application of ferrous sulfate or aluminum sulfate, which hydrolyzes to form sulfuric acid; (3) the application of a solution of sulfuric acid or of phosphoric acid; (4) the mixing of very acid soil, usually peat, with the soil. The use of sulfur is probably the most common procedure.

**Buffering in Soils.** An acid or alkaline solution or system, like a soil or plant which requires more base or acid to neutralize it than its pH value indicates, is said to be buffered. Such a system resists a change in reaction, and hence the buffer capacity of a system is often designated as its capacity to resist change in reaction. Buffering comes about through the low ionization of weak acids, weak bases, and also salts, which give rise to weak acids or bases when strong acids or bases are added. For example, a solution of phosphoric acid ( $H_3PO_4$ ), a rather weak acid, will require more base to neutralize it than would be calculated from its pH value because only a part of its hydrogen is ionized at any one time.

Soils may be buffered by the presence of carbonates, phosphates, and other salts, but generally the buffering capacity is due largely to the

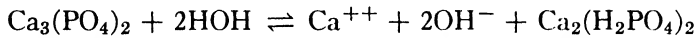
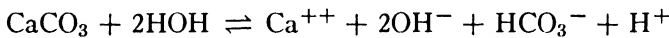
quantity of organic and inorganic colloidal material they contain. The colloidal soil complex functions as a slightly ionized acid or a slightly ionized salt of a weak acid. If the acid complex is represented by  $H_{100}X$ , the calcium salt might be written  $Ca_{50}X$ . It would be assumed, therefore, that the addition of a small amount of acid, let us say  $5H_2SO_4$ , would give a marked decrease in  $pH$ . This result would not be attained, however, because we would now have present a partially neutralized weak acid ( $Ca_{45}H_{10}X$ ) which still contains a large excess of calcium. On the other hand, the buffering effect of a strongly acid soil to the addition of alkaline material may be illustrated by the reaction



Here, as the dissociated hydrogen is neutralized, more will dissociate from the soil complex or weak soil acids, and hence little change in reaction will result.

As the buffering material of soils is usually present primarily in colloidal form, it is to be expected that soils high in clay or organic matter content will be highly buffered and hence will require comparatively large quantities of lime to neutralize them if they are strongly acid. Coarse-textured soils, on the other hand, are usually only slightly buffered.

When a soil contains an appreciable quantity of salts of weak acids, such as the carbonates, bicarbonates, phosphates, or silicates of Ca, Mg, Na, and K, they contribute to the alkalinity by hydrolysis, as shown in the reactions



If a small addition of acid is made to such a soil, the OH ions neutralized will immediately be replaced through further hydrolysis of the salt, for only a small part of the salt will dissociate at any time. It can be seen then that the buffering capacity of the colloidal complex of a soil may be greatly supplemented by the action of salts. This situation prevails in many desert soils.

**Methods of Determining Acidity.** If it is desired to measure the active acidity of a soil, it is necessary to determine the H-ion concentration. This is the most commonly used measure of acidity both in soils and in many other materials. The most accurate procedure is to use an electrode which sets up a potential with the H ions just as a copper electrode will set up a potential with the Cu ions in a solution

of CuSO<sub>4</sub>. The potential may be measured by balancing it against a half cell of known potential, making use of a potentiometer. As the potential established is proportional to the concentration of H ions, this procedure offers the opportunity for measuring the active acidity. Various electrodes have been used, such as hydrogen gas adsorbed in finely divided platinum deposited on a platinum wire; a chemical, quinhydrone, which in water functions as a hydrogen electrode; a glass tube with the end sealed and blown out into a very thin-walled bulb or into a spear-shaped point and filled with one-tenth normal HCl. The last, known as a glass electrode, is the most commonly used. Portable and relatively inexpensive sets utilizing the glass electrode are now available.

Various dyes which give distinct color shades at different pH values are used extensively, in the laboratory and especially in field kits, for determining pH values with relative accuracy. The soil may be placed either in a piece of paraffined paper, folded to make a trough, or in a depression in a porcelain plate, and the dye solution allowed to soak slowly through it. A drop of the clear percolate is drawn to the side for comparison with a standard color chart. Different dyes which change color within different pH ranges may be used separately, or several of them may be included in one solution.

A test based on the occurrence of soluble iron in acid soils involves the use of a solution of potassium sulfocyanate (KSCN). This reagent gives a red color with ferric iron.

Some tests, formerly used much more than at present, involved the determination of active plus a portion of the reserve acidity. The soil was treated with a solution of a salt, such as KCl or Ca(C<sub>2</sub>H<sub>3</sub>O<sub>2</sub>)<sub>2</sub>, the basic ion from which replaced part of the H on the acidoid with the formation of HCl or HC<sub>2</sub>H<sub>3</sub>O<sub>2</sub>. The acidity of the resulting solution was then determined by titration with a weak alkali solution or by other appropriate means.

**Interpretation of Results of Acidity Tests.** Because H-ion concentration determinations do not give a measure of total acidity, they must be used in association with other information in estimating the quantity of lime to apply in order to bring a given soil to a desired pH. In some states the actual quantity of lime needed to raise each of the principal soil types one or more pH units has been determined, and these data are used in making lime recommendations from pH determinations. In other states it is assumed that the texture of the soil correlates roughly with the cation-exchange capacity. The quantity of lime recommended for a soil with a given pH is therefore increased pro-

gressively as the soils vary from sandy soils to loams, silt loams, clay loams, and so on. Lime recommendations are usually made by experienced workers who have in mind the results of numerous field experiments involving the use of different quantities of lime on different soil types.

Several factors aside from *pH* values should be taken into consideration in deciding on the quantity of lime to apply to a given soil. For a discussion of them refer to Chapter 7.

#### CONDITIONS IN ACID SOILS WHICH ARE BENEFICIAL OR DETRIMENTAL TO THE GROWTH OF PLANTS

As stated in the introduction to this chapter, soil acidity is an indication of chemical conditions existing in a soil. The components which contribute to this situation unquestionably vary in soils of different chemical composition, and hence the same plant may not be similarly affected in all soils showing the same *pH*. This point is well illustrated by the relative need for lime of organic and mineral soils occurring in the same locality. Whereas a mineral soil with a *pH* of 5.5 may need two or more tons of limestone per acre for the growing of acid-sensitive crops, much less or possibly no lime would be needed on the organic soil. Furthermore, a scarcity of certain nutrients, as boron, in available form may occur in some soils with a high *pH*, whereas in other neutral or alkaline soils of the arid regions an excess of boron may be present.

In the following paragraphs are discussed the points raised by the list of questions concerning the conditions in acid soils which affect plant growth. This material is also presented in Chart 3.

#### Questions

1. How does the concentration of H ions in acid soil affect plant growth?
2. Are soluble iron and aluminum salts detrimental to plants?
3. How does soil acidity affect the availability of phosphorus to plants?
4. Do plants get sufficient calcium in soils of low *pH*?
5. Does soil acidity influence the activity of microorganisms?
6. May plants be classified on the basis of their tolerance of acidity?

**Effects of H Ions on Plants.** Many studies have been made to determine the influence of H-ion concentration on the growth of various plants. The results show in general that, in a nutrient solution with a concentration of H ions corresponding to that of strongly acid soils, plants grow much better than in the acid soil itself. In other words, when separated from the accompanying conditions, which pertain in an

acid soil, the H-ion concentration ceases to be especially harmful to many plants. Plants vary in this respect. It becomes necessary, then, to look for some cause in addition to H-ion concentration for the detrimental effect of acid soils on plant growth.

**Soluble Iron and Aluminum.** The quantity of soluble iron and aluminum in many soils increases as soil acidity increases. Although ferrous iron is rather toxic to many plants, ferric iron is not nearly so detrimental. Soluble aluminum, on the other hand, has a high toxicity for many commonly grown crops. In some soils the soluble aluminum present has been shown to be responsible for the larger part of the toxicity to certain crops. It is quite generally believed that soluble aluminum is one of the major factors contributing to poor plant growth in strongly acid soils.

**Phosphorus Availability in Acid Soils.** The chemical elements iron and aluminum both form compounds of low solubility with a phosphate ion in acid soils. From about *pH* 5.0 up to the neutral point, soil phosphates have an appreciable solubility on account of the basic ions present that maintain the higher *pH* and at the same time form some calcium and magnesium phosphates in the soil. In soils from about *pH* 5.0 downward, however, complex phosphates of iron and aluminum may be formed which have a very low solubility and do not supply sufficient phosphorus for plants. The calcium and magnesium phosphates are more soluble in the presence of  $\text{CO}_2$ , which comes from decaying organic matter, but this solubility effect decreases up to about *pH* 8.0. Under strongly acid soil conditions, therefore, when soluble phosphate fertilizers are applied to the soil they revert strongly to the insoluble forms, and hence on these soils the efficiency of applied phosphates is greatly reduced. Applications of lime raise the *pH* of the acid soils and tend to increase the solubility of the soil phosphorus especially when  $\text{CO}_2$  is present in the soil. Likewise, the plowing under of green manures and barnyard manures increases the  $\text{CO}_2$  content of the soil when they decay and thus helps to make the soil phosphorus available to plants.

Phosphoric acid forms three common salts with calcium, namely, monocalcium phosphate,  $\text{Ca}(\text{H}_2\text{PO}_4)_2$ , dicalcium phosphate,  $\text{CaHPO}_4$ , and tricalcium phosphate,  $\text{Ca}_3(\text{PO}_4)_2$ . The monocalcium phosphate is water-soluble, but the other two calcium phosphates are of rather low solubility. Magnesium and phosphoric acid form a similar series of phosphates, and the corresponding magnesium phosphates are more soluble than the calcium phosphates. Although the solubility of the

dicalcium and dimagnesium phosphates in water is comparatively low, these salts furnish phosphorus to plants quite readily. The tricalcium and trimagnesium phosphates, however, have only a low degree of solubility. When the soluble monocalcium phosphate fertilizer is applied to soils having a *pH* representing a medium acid to neutral condition, a large portion of the phosphorus is reverted to the more insoluble forms, but some of the phosphorus is still held in a form available to plants. It should be remembered, also, however, that plant roots intimately contact the colloidal material of the soil and thus are able to obtain more phosphorus than that represented by the immediate water-solubility.

Low solubility of phosphorus in alkaline soils results when the  $\text{Ca}_3(\text{PO}_4)_2$  hydrolyzes in the presence of  $\text{CaCO}_3$ .



The high concentration of Ca ions in the products of the reaction greatly depresses the solubility of the phosphorus. According to McGeorge and Breazeale,<sup>1</sup> this reaction takes place at a *pH* of 8.0 to 8.5. Through this *pH* range and, also somewhat below it therefore, the soil phosphorus may be highly insoluble and unavailable. At still higher *pH* ranges where the effects of the sodium and potassium ions may be appreciable, phosphorus is again held in the soil in a more soluble condition. The whole subject of phosphorus solubility in soils, however, is very complex, and it is not always easy to predict how much phosphorus will be available to plants under any set of soil conditions.

**Calcium Supply in Acid Soils.** As soils become acid only when the supply of basic ions, principally Ca, on the colloidal complex has become depleted, there is a basis for the opinion that a calcium deficiency is one cause for the restricted growth of many plants on highly acid soils. Aside from the calcium used as a nutrient, the element plays several important roles in plant nutrition, as discussed in Chapter 7. It has been found that, when calcium in a form unable to correct acidity, such as  $\text{CaCl}_2$  or  $\text{CaSO}_4$ , is supplied, the detrimental effects of acid soils on certain plants are largely overcome. There is probably considerable variation in the response of different plants to a calcium deficiency and to the application of neutral calcium salts. Furthermore, the quantity

<sup>1</sup> W. T. McGeorge and J. F. Breazeale, "The Relation of Phosphate Availability, Soil Permeability, and Carbon Dioxide to the Fertility of Calcareous Soils," *Arizona Agr. Exp. Sta. Tech. Bull.* 36, 1931, p. 375.

of exchangeable calcium in different soils having the same *pH* is decidedly variable because of differences in cation-exchange capacity. Nevertheless, a deficiency of calcium is undoubtedly a highly important factor in the growth of plants on many acid soils.

**Activity of Microorganisms.** The microorganisms which convert ammonia into nitrite and nitrite into nitrate are sensitive to acidity. The production of nitrates in strongly acid soils is accounted for, in part, at least, by the presence of decaying particles of minerals which are liberating bases, thus establishing small areas of slightly acid to neutral reaction. As the *pH* value of a soil is the summation of the reaction of all parts, there may well be innumerable very small zones of neutrality in a soil of low *pH*. One group of bacteria (*Azotobacter*) which utilizes atmospheric nitrogen without association with a legume does not function appreciably below a *pH* of 6.0. Symbiotic bacteria which grow in conjunction with several of the most widely grown legumes, such as alfalfa, sweet clover, red clover, and to a lesser extent alsike clover and soybeans, are affected adversely by moderate to strong acidity. There is considerable range in the tolerance of symbiotic bacteria to acidity. Those forming nodules on the roots of lespedeza, crimson clover, lupines, and peanuts function in soils of relatively high acidity. This question is discussed more fully in Chapter 9.

**Variation in Tolerance of Plants for Acidity.** The distribution of plants under natural conditions may well be considered due to their ability to survive under those conditions rather than an indication that the environment is the one in which they grow to best advantage. In other words, the existence of plants in a given environment is one of tolerance rather than choice. This fact is well illustrated by the rapid growth of some desert plants, such as mesquite, in a warm humid climate. Also, some acid-tolerant plants like sorrel grow much more luxuriantly in a soil which has been limed. The intensity of acidity which plants will tolerate is also influenced by the supply of available nutrients and of moisture. When other conditions are favorable for the growth of a plant it will thrive in a much more acid soil than it would if the nutrient or moisture supply were limited. Experimental studies and observation have established the fact that most crops will grow more satisfactorily in a soil which is slightly acid than in one which is neutral or alkaline. This situation pertains to acid-sensitive crops such as alfalfa.

There are some plants which grow luxuriantly only in strongly acid soils. No satisfactory explanation has been offered for this phenomenon.

In Chart 4 a number of plants are listed on the basis of their tolerance of or requirement for an acid growth medium.

No satisfactory explanation for the variation in tolerance of acidity exhibited by different plants has been advanced. A useful exercise for the student of soils is to devise several rotations, using crops which are tolerant to medium or strong acidity. These rotations should meet all the requirements of a practical rotation so far as possible.

**Chart 4. Tolerance of Plants for Acidity**

| Slightly<br>Tolerant | Moderately<br>Tolerant | Highly<br>Tolerant | Very Highly<br>Tolerant * |
|----------------------|------------------------|--------------------|---------------------------|
| Alfalfa              | Austrian winter pea    | Crotalaria         | Azalea                    |
| Ladino clover        | Crimson clover         | Lupine             | Rhododendron              |
| Red clover           | Hop clover             | Kudzu              | Blueberry                 |
| Sweet clover         | Lespedeza              | Velvet bean        | Watermelon                |
| Sugar beets          | Alsike clover          | Corn               | Cranberry                 |
| Beans                | Vetch                  | Oat                |                           |
| Table beets          | Cowpea                 | Rye                |                           |
| Asparagus            | Cotton                 | Buckwheat          |                           |
| Cabbage              | Peanut                 | Sorghum            |                           |
| Cauliflower          | Barley                 | Millet             |                           |
| Spinach              | Soybean                | Redtop             |                           |
|                      | Wheat                  | Tobacco            |                           |
|                      | Cantaloupe             | Strawberry         |                           |
|                      |                        | Peach              |                           |
|                      |                        | Pecan              |                           |

\* Are harmed by lime except under exceptional conditions.

#### CONDITIONS OF DEVELOPMENT AND EFFECT OF NEUTRAL AND ALKALINE SOILS ON PLANTS

A large proportion of the soils in the Chernozem belt and westward has a neutral or an alkaline reaction. Exceptions are found in the soils occupying areas of high rainfall near the coast and in certain intermountain valleys. Some desert soils have a *pH* of slightly below 7. East of the Chernozems many areas of soil are non-acid because they have one or more of the following characteristics: (1) location in which seepage water passing through lime-carrying soil material supplies lime to the soil; (2) young soils derived from limestone or lime-rich material and from which the Ca has not yet been leached; (3) soils developing on recently exposed lake beds or on poorly drained soil material containing  $\text{CaCO}_3$ ; (4) lateritic soils in which the decay of organic matter is sufficiently rapid to maintain a supply of bases.

Neutral and alkaline soils occupy approximately half of the area of continental United States, and hence they deserve much consideration.

The discussion of this important group of soils will be divided into sections, as indicated by the following questions.

### Questions

1. Does limited rainfall contribute to the neutrality and alkalinity of soils?
2. Why are "black alkali" soils so named?
3. How are "white alkali" soils developed?
4. What plants are tolerant of alkali in soils?
5. In what ways are high salt concentrations detrimental to plants?
6. What methods are used to improve alkali soils?

**Low Rainfall and Soil Reaction.** It has been shown previously that soluble salts, principally carbonates and bicarbonates of Ca, Mg, Na, and K, are produced during the decomposition of rocks and minerals. When rainfall is sufficient to leach from the soil the Na and K salts but only a part of the Ca and Mg carbonates, the latter accumulate and keep the colloidal complex saturated. Hence a neutral or slightly alkaline reaction is maintained. With less rainfall a copious accumulation of Ca and Mg carbonates occurs and a decidedly alkaline reaction develops. When conditions permit some of the  $\text{Na}_2\text{CO}_3$  to remain in the soil, a pH of 8.5 or higher may be attained and the term alkali is then usually applied.

**Black Alkali.** When  $\text{Na}_2\text{CO}_3$  occurs in a soil it readily hydrolyzes, forming NaOH:  $\text{Na}_2\text{CO}_3 + 2\text{HOH} = 2\text{NaOH} + \text{H}_2\text{O} + \text{CO}_2$ . This strong base attacks organic material to form a dark-colored soil solution which may result in a black or brownish black residue on the surface of the soil. As a result the term black alkali has been applied to such soils. Not only is the alkali corrosive to plant tissue, but also the Na is adsorbed on the colloidal complex and causes a puddled soil structure which greatly impedes the development of roots and the absorption of irrigation water.

**Saline Soils.** In some desert soils an accumulation of the sulfates of Ca, Mg, and Na, and sometimes of NaCl and KCl, accompanies the carbonates of Ca, Mg, and Na. Such soils have a light gray color and are given the name of white alkali. The source of the soluble salts in saline soils is sometimes difficult to explain. In some the problem is simplified through evidence that the area is the bed of an extinct saline lake or of an area of the ocean. There is no such evidence in many soils, however. Seepage water passing through rocks impregnated with salts is sometimes the explanation. The evaporation, over long periods, of drainage water containing small quantities of salts culminates in appreciable salt accumulations in some basins.

**Plant Growth in Neutral and Alkaline Soils.** Plants vary in their tolerance of alkalinity and salt content of soils just as they do in their tolerance of acidity. In Chart 5 are listed a number of crops and native plants which are especially tolerant of alkalinity and also several which are less tolerant.

**Chart 5. The Relative Tolerance of Some Western Plants for Salt Constituents in the Soil Solution**

| Strong Tolerance     | Good Tolerance *         | Medium Tolerance *  | Weak Tolerance      |
|----------------------|--------------------------|---------------------|---------------------|
| Sugar beets          | Barley (hay crop)        | Onions              | Beans, wax          |
| Garden beets         | Tomatoes                 | Squash              | Beans, navy         |
| Milo                 | Alfalfa                  | Carrots             | Field peas          |
| Bermuda grass        | Cotton                   | Asparagus           | Horsebean           |
| Rhodes grass         | Sorgo                    | Cowpeas             | Red clover          |
| Alkali sacaton grass | Kale                     | Rice                | Vetch               |
| Rescue grass         | Rape                     | Rye (grain crop)    | Proso               |
| Greasewood           | Perennial ryegrass       | Barley (grain crop) | Oats (grain crop)   |
| Seepweed             | Wild ryegrass            | Wheat               | Emmer (grain crop)  |
| Pickleweed           | Italian ryegrass         | Oats (hay crop)     | Peaches, Elberta on |
| Saltgrass            | Crested wheatgrass       | Grain sorghums      | Lovell rootstock    |
| Salt sage            | Slender wheatgrass       | Foxtail millet      | Sage brush          |
| Tussock-grass        | Mountain brome-<br>grass | Flax                | Creosotebush        |
| Date palms           | Tall fescue              | Guayule             | Mesquite            |
|                      | Reed canary grass        | Sunflower           | Little rabbitbrush  |
|                      | Sudan grass              | Ladino clover       | Gallentia           |
|                      | Blue grama grass         | Strawberry clover   | Giant wild-rye      |
|                      | Desert saltbush          | Sweet clover        | Corn                |
|                      | Shadscale                | Smooth bromegrass   | Potatoes            |
|                      | Winterfat                | Meadow fescue       | Oranges             |
|                      |                          | Orchard grass       | Pears               |
|                      |                          | Dallis grass        | Figs                |
|                      |                          | Tall oatgrass       | Apples              |

\* A few crop plants listed in the medium and good groups do not die in strong salt concentrations, but they fail to make appreciable growth.

Black alkali is harmful to the tissues of living plants, since it attacks them in the same way that it does the organic matter of the soil.

A concentration of soluble salts, such as occurs in white alkali soils, is not corrosive to plant tissue but may prevent the absorption of moisture by the plants because of high osmotic pressure. Likewise, the presence of some of the ions in considerable concentration may limit the intake of certain nutrients, as experiments have shown that the absorption of a given nutrient is influenced by the presence or absence of other ions in the nutrient solution. There are data which indicate

that negatively charged nutrient ions are absorbed with difficulty by plants when the soil reaction exceeds pH 7.6. The use of a strongly acid fertilizer has been found to facilitate the intake of nutrients from strongly alkaline soils.

The solubility of the compounds of some elements which are required by plants is also suppressed by alkalinity. The insolubility of phosphates in a soil containing calcium carbonate is an example. Manganese is also usually found in a rather insoluble condition in alkaline soils, as is also iron. On the other hand, the salts of certain elements may be present in sufficient concentration to be toxic to planted crops.

**Correction of Alkali.** Under irrigation the alkalinity of soil may increase, owing to the upward movement of salts with capillary water. Several methods of treating soil to counteract the detrimental effects of black alkali have given more or less satisfaction. Ground sulfur applied to the soil oxidizes to sulfuric acid and converts the  $\text{Na}_2\text{CO}_3$  to  $\text{Na}_2\text{SO}_4$  and  $\text{H}_2\text{CO}_3$ . The  $\text{H}_2\text{CO}_3$  so liberated combines with  $\text{CaCO}_3$  to make  $\text{Ca}(\text{HCO}_3)_2$ , which tends to develop a granular structure in the soil. Sulfuric acid and  $\text{CaSO}_4$  produce the same results, and both are used although  $\text{CaSO}_4$  is used much more widely. Decaying organic matter has a very salutary effect, since the  $\text{CO}_2$  produced converts the calcium into the relatively soluble bicarbonate, in which form it is effective in inducing granulation. It is considered good practice to apply manure or to plow under a green-manuring crop in conjunction with the application of ground  $\text{CaSO}_4$ . Regardless of the soil corrective used in reclaiming alkali soils, the products of the reaction between the corrective and the alkali must be subsequently removed by leaching with irrigation water. An occasional heavy application of irrigation water is good practice because it tends to prevent the accumulation of alkali salts. The subject of the management of alkali soils is discussed more fully in Chapter 18.

## 7

## Lime and Its Use

Liming is an important practice in scientific soil management. Too frequently soil fertility is regarded chiefly as a matter of supply of nitrogen, phosphorus, and potassium, and insufficient emphasis is placed on calcium, which performs many indispensable functions in plant growth and in modifying soil conditions. The liming of acid soils should be accepted by all as a fundamental and essential practice; it is frequently referred to as the "backbone" of permanent agriculture in humid regions.

In the discussion of soil acidity, it was pointed out that unfavorable soil conditions associated with acidity may be corrected by the application of lime. When the chemist speaks of lime, he usually refers to calcium oxide ( $\text{CaO}$ ), but from the agricultural point of view a more general meaning is implied by the term. Agriculturally, lime may be said to be any compound of calcium or of calcium and magnesium *capable of counteracting the harmful effects of an acid soil.*

It is obvious why compounds of calcium and magnesium are used for liming purposes rather than compounds of such elements as sodium and potassium. In humid regions the ions of calcium and magnesium are the usual predominating cations in soils, and it is natural that an attempt should be made to restore the supply when depleted. Moreover, materials carrying calcium and magnesium are relatively abundant, widespread, and cheap. These materials are effective in neutralizing all types of soil acids and have no detrimental effects on the physical properties of soils and are not toxic to plants in amounts normally applied. Potassium salts are too expensive to use for correcting acidity, and alkaline salts of either potassium or sodium encourage the development of a poor physical condition of the soil. Furthermore, these salts are caustic to handle and are toxic to plants if applied in large amounts.

Liming materials are used to bring about chemical, physical, and biological changes in the soil which are beneficial to plant growth. If used intelligently they give satisfactory results, but if unwisely used

they may prove detrimental to soils and crops. This subject of lime and its uses will be discussed under the following topics.

### Objectives

- A. The need of soils for lime.
- B. Functions of lime in the soil.
- C. Forms of lime used for application to the soil.
- D. Guarantees of chemical composition and fineness of lime.
- E. Sources of lime for agricultural use.
- F. Use of lime on acid soil.

### THE NEED OF SOILS FOR LIME

It has been estimated that perhaps 70 per cent of the tillable land in the eastern half of the United States is acid and requires liming for maximum crop production. A large part of this area at one time contained sufficient calcium for optimum growth of legumes and other lime-loving crops. But this calcium supply has greatly diminished, and in most of eastern United States the supply is no longer sufficient. Certain areas on the West Coast, where there is considerable rainfall, are also acid. Liming has been practiced in some of the eastern states since colonial days, and the use of lime has gradually spread westward, even into regions of limited rainfall. There are several important reasons why the popularity of lime as a soil fertilizer and soil amendment is so widespread.

### Questions

1. Were the parent materials of all soils well supplied with lime?
2. Does lime readily leach from soils?
3. Are large quantities of lime removed from soils by crops?
4. Do the acid substances normally added to soils tend to increase the need of soils for lime?

**Soils Developed from Acid Rocks.** Certain soils have developed from rocks and minerals which in themselves were naturally acid. Those rocks which contain more quartz or silica than basic constituents are known as acid rocks. Soils developed from such materials would be acid even though no loss of bases occurred during soil formation (see p. 116). There are considerable areas of siliceous sandy soils which have always been in need of lime. Most acid soils, however, have developed as a result of leaching losses and crop removal of bases.

**Need for Lime Increased by Leaching.** As has been indicated, the amount of rainfall and the lime needs of soils are closely related.

No one factor tends to remove the basic materials from the soil in humid regions to so great an extent as leaching, and calcium is the element which usually suffers the greatest loss. As a rule, the basic elements in soils are more readily soluble than the acid constituents and hence are more readily carried away in drainage water as it percolates through the soil. With a continuation of this process, it is obvious that a deficiency of bases and the development of acid soils will occur sooner or later. Since calcium and, to a lesser extent, magnesium are readily lost from soils and since calcium and magnesium are the active constituents of lime, soils which do not supply them in sufficient quantities for plant growth are designated as lime-deficient.

The leaching losses of calcium and magnesium from soils depend to a large extent on the quantities of these elements present and the forms in which they are held, the amount of water percolating through the soil, the vegetative cover, and the production of carbonic acid. Calcium and magnesium for the most part are held in the soil in either the silicate or the carbonate form or as exchangeable ions on the soil colloids, and both forms are subject to leaching losses. Owing to the continuous production of carbon dioxide through organic matter decomposition and plant excretion and to the consequent development of carbonic acid, there is a constant tendency toward a replacement of calcium and magnesium ions on the soil colloids by the hydrogen ions of the carbonic acid. These basic ions are thus released and are susceptible to loss by leaching. Furthermore, the carbonates of calcium and magnesium in the soil may be gradually converted to the more soluble bicarbonates. Leaching losses are much greater from fallow fields than from comparable fields growing crops. This difference is due to the smaller quantities of percolating water in cropped land and to the intake of bases by the plants.

**Crop Removal of Bases.** The removal of bases through cropping is greater than the removal of acids; thus the soil is left more acid. Such crops as the legumes, particularly alfalfa and clover, require considerably more calcium and magnesium than any other crops and consequently make a heavy demand for bases. An average crop of alfalfa will remove about 100 pounds of calcium per acre; clover, about 60 pounds; and soybeans, 30 pounds. Cropping, therefore, tends to deplete the lime content of the soil (see p. 314).

**Acid Substances that Increase the Need for Lime.** Near industrial areas large quantities of sulfur are carried into the soil with the precipitation. This sulfur is readily changed to an acid and as such encourages the loss of soil bases. Decaying organic matter also pro-

duces various acids, and the process of nitrification results in the formation of nitrates which greatly facilitate the loss of bases. These acids produce salts which on the whole are quite soluble. The use of certain fertilizers (see p. 351) is known to increase the need for lime.

The need of soils for lime, then, is caused by the loss of the basic constituents, calcium and magnesium, through leaching and cropping and is aggravated by other processes or practices which add acid constituents to the soil, although, as pointed out, a need for lime may exist even though no loss of bases occurs during soil formation.

Lime was applied as a soil amendment in ancient times, long before the Christian Era, and the same factors which created lime-deficient soils at that time are still operating today. Soils in humid areas will always be affected by these lime-depleting processes.

### FUNCTIONS OF LIME IN THE SOIL

Lime produces several specific effects, which result in the improvement of soils and increased crop production. Since soils may vary greatly in their physical, chemical, and biological properties, it is not to be expected that a lime deficiency will produce exactly the same unsatisfactory conditions in each case, and, as a result, some soils may give a much greater response to lime than do other soils. Injurious effects may result from the use of lime under certain conditions, especially when applied in excess. Lime has several important functions in the soil, as indicated by the questions which follow.

#### Questions

1. Are all forms of calcium effective in neutralizing soil acids?
2. Does the calcium ion have a direct effect on plant growth?
3. What is the effect of liming on the solubility of soil mineral elements?
4. Are the desirable microbiological soil processes favorably influenced by the presence of lime?
5. Does the presence of lime affect the chemical nature of soil colloids?
6. What are the effects of lime on soil structure?
7. Does the application of lime increase the effectiveness of fertilizers and manure?

**Neutralization of Soil Acids.** Since the accumulation of hydrogen ions in soils is accompanied by the loss of basic elements, chiefly calcium, an addition of lime will correct the calcium deficiency and at the same time neutralize soil acidity. Acid soils are beneficially changed in character by the application of calcium compounds in the oxide,

hydroxide, or carbonate form but not by the addition of calcium in the sulfate (gypsum) or chloride form. When the last two compounds are added to acid soils, strong, stable mineral acids ( $H_2SO_4$  and  $HCl$ ) are formed, and unless they are leached from the soil no decrease in acidity can result although the quantities of soluble calcium in the soil may be increased by the application of these substances.

Lime in the forms of oxide, hydroxide, and carbonate reacts rather rapidly with moist, acid soils. The reactions of these forms vary somewhat, but the final result is the same. When lime is added, the soil water becomes charged with calcium ions which have the ability to replace (by ionic exchange) the hydrogen ions on the colloidal complex. The hydrogen ions are released, and they unite with hydroxyl ions to form water. When the colloidal clay becomes saturated with calcium, it is no longer acid; the acidity of the soil has been neutralized or corrected. In other words, the process of neutralizing acid soils is the reverse of that by which they become acid, as explained in Chapter 6 (p. 117).

**Influence of Calcium on Plant Growth.** Calcium is an indispensable element for the growth of all crops. Many soils are too low in lime content to supply the needs of crops, especially certain species of leguminous plants. All the functions calcium performs within the plant are not definitely known, but it is generally believed that calcium has an influence on the translocation of carbohydrates and certain mineral elements within the plant and on the development of roots. It appears to be essential for cell-wall construction and perhaps aids in neutralizing organic acids within the plant or in regulating the acid-base balance in plants. Calcium may also influence, either favorably or unfavorably, the absorption of other elements. Within certain limits, for example, an inverse relationship has been found between the intake of calcium and potassium by plants. It may counteract to some extent the toxic effects of high concentrations of magnesium and sodium.

The nutritive value of animal feeds is to a large extent influenced by the calcium content of plants. It is important in the development of the bones of animals and of the shell of eggs. That lime should be present in sufficient quantities in all properly balanced livestock rations is now an accepted fact. The importance of the calcium content of food for growing children and for treating certain types of human disease is well recognized by physicians.

**Effect of Lime on the Solubility of Soil Mineral Elements.** As a general rule, a relatively high percentage of the phosphorus in soils well supplied with lime is available for plant use, and the phosphorus

of calcium-deficient acid soils is in relatively unavailable forms even though the total phosphate content is comparatively high. Soil phosphorus is generally most readily available to plants in neutral or slightly acid soils, and with increasing acidity its availability decreases. In strongly acid soils, in the presence of aluminum and iron compounds, soluble phosphates combine with these elements, forming relatively insoluble aluminum and iron phosphate compounds. In alkaline soils in the presence of excess calcium carbonate (about 2 per cent), phosphorus combines with calcium, forming tricalcium phosphate. This form of phosphorus is of low solubility but is more soluble than the phosphates of aluminum and iron.

Liming acid soils has a tendency to make the phosphorus more available by converting a part of it that is present as aluminum and iron phosphate to the more available calcium phosphates. The more desirable forms from the standpoint of plant use are the monocalcium and dicalcium phosphates. It is also likely that liming results in the liberation of the organic phosphorus in the soil through stimulation of decomposition processes. It is obvious, however, that lime alone will not solve the problem of phosphorus availability because many soils are so depleted of phosphorus that lime has little effect in increasing crop yields unless accompanied by applications of phosphate fertilizer.

The effect of lime on the solubility of soil potassium cannot be stated with any degree of certainty. Theoretically, calcium by the process of ionic exchange should displace or liberate some of the exchangeable potassium of the soil. Under field conditions this is difficult if not impossible to demonstrate because of so many complicating chemical and biological factors. There is experimental evidence indicating that the solubility of potash is decreased in certain soils that have been overlimed. Marked crop response to potash fertilizers is often secured on such soils. It is perhaps safe to say that lime does not greatly influence the availability of potassium.

When soils become deficient in bases, the solubility of aluminum, iron, and manganese increases; in strongly acid soils, the high concentrations of these elements may be toxic to crop plants. Excess quantities of these elements may become available at *pH* values below 5.5. Soils having reactions between *pH* 5.5 and *pH* 7.0 usually supply plants with sufficient quantities of both iron and manganese, but at *pH* values above 6.5 or 7.0, especially in sandy soils, they may become insoluble to such an extent that plants are unable to satisfy their needs. Caution is therefore necessary in the use of lime. A normal application may

prevent toxic concentrations of these substances; yet too much lime may create an iron and manganese deficiency.

**Effect of Lime on Microbiological Processes of the Soil.** Calcium is closely associated with certain important microbiological processes. The more important effects of calcium on the soil population may be grouped as follows: (1) promoting the decomposition of organic matter; (2) making conditions favorable for nitrification and sulfofication; and (3) providing favorable conditions for the growth and functioning of both symbiotic and non-symbiotic nitrogen-fixing bacteria. Each group of organisms which contributes to the above processes functions most efficiently in soils well supplied with lime. In many of these processes, it is not necessarily a matter of changing the *pH* but one of supplying soluble calcium.

Lime may be used as a preventive of certain types of plant diseases which occur only in acid soils. However, an alkaline soil, caused by the use of too much lime, is likely to depress certain desirable microbiological processes, such as nitrification and the decomposition of organic matter. Furthermore, it is generally believed, although it does not always occur, that the application of lime in amounts sufficient to make a soil neutral or alkaline favors potato-scab disease.

**Effect of Lime on the Chemical Nature of Soil Colloids.** As has been explained, acid substances of soils are, for the most part, not soluble in the ordinary sense, but rather they are colloidal. Furthermore, calcium is usually the dominant exchangeable cation in neutral soils in humid regions. Under leaching conditions the calcium and other bases are gradually replaced by hydrogen ions from the surface of colloidal particles. As these particles become more acid, they tend to become more chemically unstable. Silica splits off from the inorganic colloidal materials and their cation-exchange capacity is greatly reduced. The maintenance of a high cation-exchange capacity in soils is thus mainly dependent on the maintenance of a high calcium content. Aluminum may also come into play as an exchangeable cation. The remedy for this is lime. Not merely does it act as a neutralizing agent, but also it restores calcium on the soil particles, producing significant effects on various chemical as well as physical properties of the soil.

**Effect of Lime on Soil Structure.** For many years it was thought that the presence of sufficient calcium in the soil to replace most of the hydrogen on the colloids would result in granulation and a desirable soil structure. The improvement of soil structure was one of the benefits promised from liming an acid soil. More recently careful studies have shown that replacement of hydrogen by calcium on the soil colloids does

not necessarily result in a granular condition of the soil. In fact, soils high in lime content are known which have a dense structure almost impervious to water. The presence of much calcium on the colloidal complex does lead to flocculation of the colloids, but this fact does not insure a granular soil structure. It seems, then, that the effect of lime upon granulation may be indirect; that is, it may come about through the influence of lime on the production and decomposition of organic matter.

**Influence of Lime on the Effectiveness of Fertilizers and Manure.** Fertilizers and manures are more effective when all soil conditions are favorable. Seldom do they produce maximum results in sour soils. Lime is not a substitute for fertilizers and manure, but when lime is needed its application will greatly increase their effectiveness.

In soils deficient in lime, various substances, like iron and aluminum compounds, are in a condition to combine with soluble phosphorus, applied in fertilizers and manure, and change it into less soluble forms. Since the solubility of the natural phosphorus compounds is also low in lime-deficient soils, it is evident that crops are likely to suffer from the need of phosphorus in acid soils.

Nitrogen and potassium of fertilizers and manure are not made insoluble as a result of a lime deficiency, but, since crops do not grow so well as in lime-rich soils, the plants cannot utilize the nitrogen and potassium as well as they could if growing conditions were made more favorable by the presence of lime. If nitrogen is applied in forms other than nitrate, the presence of lime will hasten the production of nitrates.

### FORMS OF LIME

Lime may be purchased in three chemical forms, namely, the oxide, the hydroxide, and the carbonate. They all have the advantage of leaving no harmful residue in the soil. The oxide and hydroxide change to the carbonate and bicarbonate forms in moist soils at a fairly rapid rate. The calcium and magnesium of the bicarbonate form may replace the hydrogen ions on the colloidal clay with the resultant formation of carbonic acid which readily breaks up into water and carbon dioxide. On the other hand these bicarbonates may hydrolyze to form  $\text{Ca}(\text{OH})_2$ , the Ca of which replaces the adsorbed hydrogen ions which in turn combine with the OH ions to form water.

Calcium silicate, which is a constituent of blast-furnace slags, can be classed as a liming material. It supplies calcium, neutralizes acidity, and leaves no harmful residue. It has not been used extensively in the

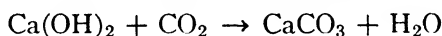
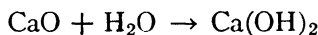
past, but in the future it may be classed as an important form of lime in areas near the points of production. In the discussion of the different forms of lime the following questions will be answered.

### Questions

1. How are the different forms of lime prepared, and what are some of their outstanding properties?
2. What is the relative neutralizing power of the different forms of lime?
3. Are the different forms of lime equally effective in the soil?

**Carbonate Forms of Lime.** Calcium carbonate is commonly referred to as carbonate of lime, under which name several lime materials are sold. Of the various chemical forms of lime, this is the most widely used for liming purposes; and most of the naturally occurring liming materials exist in the carbonate form.

By-product lime from various industries contains what is frequently referred to as precipitated carbonate. Such carbonates are usually produced from either calcium oxide, CaO, or calcium hydroxide, Ca(OH)<sub>2</sub>. Calcium carbonate, CaCO<sub>3</sub>, may be produced by combining calcium oxide and carbon dioxide, CO<sub>2</sub>. In this reaction 56 pounds of calcium oxide combine with 44 pounds of carbon dioxide to produce 100 pounds of calcium carbonate. The steps in this reaction are



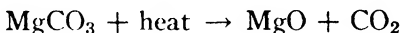
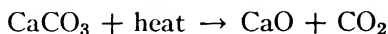
The calcium in limestones is mainly in the carbonate form. Many such stones are nearly pure calcium carbonate. Certain other limestones contain various percentages, up to about 45 per cent, of magnesium carbonate with the calcium carbonate. The latter limestones are often referred to as dolomitic or high-magnesium stones. The term high calcium is applied to limestones containing little magnesium. Any stone containing less than about 80 per cent of carbonates is usually considered low-grade, and those stones having more than 95 per cent of total calcium and magnesium carbonates are preferred for liming purposes.

Marl, chalk, marble, coral, and shells also contain lime in the carbonate form in varying proportions.

Calcium carbonate is only slightly soluble in pure water, but, if the water is saturated with carbon dioxide at ordinary temperature, it is soluble to the extent of about 1 part by weight in 1,000 parts of water, forming calcium bicarbonate, Ca(HCO<sub>3</sub>)<sub>2</sub>.

**Oxide Forms of Lime.** Calcium oxide is known under several commercial names, such as lime, burned lime, quicklime, caustic lime, stone lime, lump lime, and unslaked lime. Commercial oxide of lime is usually marketed in paper bags and in a pulverized form, although some is still sold in the granular or lump condition. The oxide form of lime is highly caustic and disagreeable to handle.

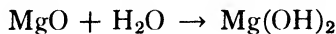
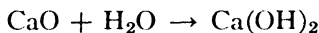
Commercial oxide of lime is prepared by heating any form of carbonate of lime,  $\text{CaCO}_3$  and  $\text{MgCO}_3$ , the carbon dioxide being driven off as a gas and the calcium oxide and magnesium oxide remaining as a solid residue.



On burning 100 pounds of pure  $\text{CaCO}_3$ , 44 pounds of  $\text{CO}_2$  are lost and 56 pounds of  $\text{CaO}$  remain. There is no loss of calcium in burning; the calcium oxide contains all the calcium of the original material.

The impurities commonly found in commercial oxide forms of lime consist of the original impurities of the limestone or other carbonate which is burned. These impurities usually consist of clay, sand, and compounds of iron and of aluminum. The oxides readily take up water and in turn carbon dioxide to form hydroxides and carbonates, respectively. Furthermore, if the burning has been incomplete (semi-calcined lime), a considerable amount of the original carbonate may remain. Although commercial oxide of lime varies greatly in composition, the ratio of calcium oxide to magnesium oxide is usually 4 or 5:1.

**Hydroxide Forms of Lime.** When calcium or magnesium oxides come in contact with water, they undergo what is known as slaking; water combines with the oxides, forming compounds known chemically as calcium hydroxide and magnesium hydroxide or commercially as slaked lime, caustic lime, hydrated lime, hydrate, agricultural hydrate, or water-slaked lime.



According to the first reaction, 56 pounds of calcium oxide combine with 18 pounds of water to form 74 pounds of calcium hydroxide.

Commercial slaked lime is thus impure calcium and magnesium hydroxides. It is a mixture of hydroxides and carbonates together with such impurities as were present in the original lime before slaking. Furthermore, if the slaking process was not carried to completion, it will

contain a corresponding amount of oxide. Thus, six important chemical forms of lime may be present, in hydrated lime, the oxides, hydroxides, and carbonates of both calcium and magnesium.

When burned lime combines with water, marked physical changes occur. The lime increases in bulk, breaks down to a very fine powder, and generates much heat. One volume of burned lime of good quality, on slaking, increases two or three times in volume.

Commercial hydrate appears on the market as a white powder and is usually handled in paper bags, although, if produced locally, bagging is not always necessary. Slaking sometimes is done in small piles in the field and the product spread directly from the piles. The hydrate is similar to the oxide forms in that it is caustic and disagreeable to handle.

Air-slaked lime is calcium oxide (together with magnesium oxide) that has absorbed moisture from the air and changed into hydroxide. As the slaked lime is formed, it slowly absorbs carbon dioxide from the air, forming carbonate (see p. 142). Air-slaked lime usually contains larger quantities of carbonate and less of hydroxide than water-slaked lime.

**Silicate Forms.** Blast-furnace (basic) slag is a by-product of pig-iron manufacture from iron ore and limestone. In the smelting process, silica of the ore reacts with calcium of the limestone to form calcium silicate. Carbon dioxide is driven off during the burning process, and there are also some magnesium and calcium oxides in the slag. It is essential that this material be finely ground in order to be effective. It is used as a liming material to a considerable extent in some of the southern states.

**Relative Neutralizing Power of the Different Forms of Lime.** The neutralizing power of liming materials is always calculated on the basis of pure calcium carbonate as 100 per cent. It has been pointed out that, when pure calcium carbonate is burned in a kiln, 100 pounds of the dry material give off 44 pounds of carbon dioxide gas, leaving 56 pounds of calcium oxide. When the 56 pounds of calcium oxide are moistened, they react with 18 pounds of water to form 74 pounds of hydrated lime. It is obvious, then, that 100 pounds of pure calcium carbonate, 74 pounds of pure calcium hydroxide, and 56 pounds of pure calcium oxide all contain the same amount of calcium and all have the same power to neutralize soil acidity because it is the calcium ions that displace the hydrogen ions in correcting acidity.

The relative ability of different liming materials to correct acidity frequently is expressed on a percentage basis. Thus, the neutralizing

power of the different forms of lime in the pure state is determined by their molecular weights. The molecular weight of calcium carbonate is 100; of calcium hydroxide, 74; and of calcium oxide, 56. By dividing 100 by 74 the figure 1.35 is obtained, which means that 1 pound of calcium hydroxide supplies the same amount of calcium as 1.35 pounds of calcium carbonate. In other words, if expressed on a percentage basis ( $1.35 \times 100$ ), pure calcium hydroxide has a neutralizing value of 135 per cent relative to calcium carbonate. Likewise, calcium oxide ( $100/56 \times 100 = 178$ ) has a neutralizing power of 178 per cent. Pure magnesium carbonate with a molecular weight of 84 has a neutralizing value of 119 per cent ( $100/84 \times 100 = 119$ ). A limestone containing 80 per cent calcium carbonate and 20 per cent magnesium carbonate would have a neutralizing value of 103.8 per cent ( $80 + 20 \times 1.19 = 103.8$ ). Frequently magnesium limestones have a neutralizing power of 107 or 108 per cent. The molecular weights, neutralizing values, and calcium carbonate equivalents for the common chemical forms of liming materials in a pure state are given in Table 9.

TABLE 9  
RELATIVE NEUTRALIZING POWER OF DIFFERENT FORMS OF LIME

| Form of Lime        | Molecular Weight | Neutralizing Value, per cent | Pounds Equivalent to 1 Ton of Pure CaCO <sub>3</sub> |
|---------------------|------------------|------------------------------|--|
| Calcium carbonate   | 100              | 100                          | 2,000  |
| Magnesium carbonate | 84               | 119                          | 1,680  |
| Calcium hydroxide   | 74               | 135                          | 1,480  |
| Magnesium hydroxide | 58               | 172                          | 1,160  |
| Calcium oxide       | 56               | 178                          | 1,120  |
| Magnesium oxide     | 40               | 250                          | 800  |

As indicated above, pure hydrated lime made from a pure calcium limestone would have a neutralizing power of about 135 per cent, but, if made from a stone high in magnesium, the neutralizing power may run as high as 150 per cent. If all the oxides of calcium or magnesium were not converted into the hydroxide form during the process of hydration, the so-called hydrated limes may have a neutralizing power of about 170 per cent. Theoretically, pure calcium oxide has a neutralizing value of 178 per cent, but, if commercial oxide of lime is made from a magnesium stone, the value may be higher if the stone is completely burned; and, if incompletely burned, leaving considerable quan-

tities of carbonate, the neutralizing value may be considerably less than the theoretical value for calcium oxide.

**Effectiveness of Different Forms of Lime.** In considering the effectiveness of a liming material, it is to be remembered that the total neutralizing power is not the only factor of importance. The density, hardness, and fineness of grinding of limestone, for example, may greatly influence the speed of its action in the soil. Dolomitic stones are harder and dissolve more slowly than calcium stones, and for this reason it is not customary to consider the higher neutralizing value of magnesium stones in determining the amount of lime to apply. In other words, a ton of a high-calcium stone seems as effective as a ton of a high-magnesium stone in correcting an acid soil.

Burned lime ( $\text{CaO}$ ) is changed to hydrated lime under ordinary moist soil conditions before it has time to act chemically with the soil; consequently, it would not be expected to be more effective than hydrated lime applied directly. Theoretically, the hydrate (and oxide) form of lime can be expected to react in the soil somewhat more rapidly than finely ground limestone, because the former is a very fine powder and can therefore be more thoroughly distributed in the soil. Moreover, freshly slaked lime consists mostly of calcium hydroxide, which is more soluble in water than is the calcium carbonate of ground limestone. This difference in solubility may result in its being more thoroughly and uniformly distributed through the soil if rain follows soon after the application of lime. Consequently, because of greater solubility and more thorough distribution in the soil, calcium hydroxide should be more active than the carbonate in producing chemical changes in the soil. It is to be emphasized, however, that calcium hydroxide will sooner or later change into the carbonate form in the soil. Under certain conditions this change may require several weeks, and during that time the hydroxide form can be considered more chemically active than the carbonate. But, over a period of years, or even a rotation, there seems to be little difference in the effectiveness of the various forms of lime when applied in equivalent quantities. Therefore, the conclusion is reached that the effectiveness of a liming material in correcting an acid soil depends on its content of calcium and magnesium and is little influenced by the particular chemical form (hydroxide, oxide, or carbonate) in which these bases occur. It is assumed that if the carbonate form is used it is ground to a satisfactory degree of fineness. Table 10 gives the relative quantities of the different forms of lime used in sections of the United States in 1948.

TABLE 10

APPROXIMATE CONSUMPTION OF LIMING MATERIALS IN DIFFERENT REGIONS  
AND IN THE UNITED STATES AS A WHOLE DURING 1948 \*

| Region                   | North-east        | East Central | North Central | Southern | Western | Total  |
|--------------------------|-------------------|--------------|---------------|----------|---------|--------|
| Forms of Lime            | Tons, 000 omitted |              |               |          |         |        |
| Agricultural limestone † | 1,826             | 2,285        | 17,312        | 1,316    | 1,025   | 23,764 |
| Burned lime              | 28                | 85           | ...           | ...      | 11      | 125    |
| Hydrated lime            | 109               | 44           | 18            | 1        | 2       | 174    |
| Marl                     | 1                 | 73           | 441           | ...      | 1       | 516    |
| Miscellaneous ‡          | 3                 | 23           | 133           | 68       | 6       | 233    |
| Totals                   | 1,967             | 2,510        | 17,904        | 1,385    | 1,045   | 24,812 |

\* Courtesy of National Lime Association.

† Includes all the various grades of limestone used for direct application; from the finely ground material (all of which passes a 20-mesh sieve) customarily used in the East, to coarser grades used in other areas. Also includes limestone used in fertilizer mixtures.

‡ Includes mollusk and egg shells, refuse from paper mills, sugar mills, and carbide plants, slag, and lead-mine chats.

### LIME GUARANTEES

Several states have laws regulating the sale of commercial lime materials. The regulations deal with both the guaranteed chemical composition and the fineness. Although there is little uniformity in the provisions of the lime-control laws of the different states, they all have as their aim the establishment of certain standards or methods of expressing guarantees that may be used for comparative or evaluation purposes. The topic is discussed under two headings, as indicated by the questions.

#### Questions

1. What is included in the chemical guarantee, and what are the various ways in which it may be expressed?
2. Of what value is the guarantee of fineness?

**The Chemical Guarantee.** Since the potential capacity of a liming material to neutralize soil acidity is determined by its content of calcium (and/or magnesium) and since these materials are sold on the basis of their neutralizing power, chemical guarantees are of great importance. The chemical guarantees for commercial oxide and commercial hydrated lime may be stated in several different ways. The guarantee may be phrased in terms of oxide content, calcium oxide equivalent, neutralizing power, or the percentages of elemental calcium and magnesium. In order to clarify the meaning of these different methods of expression, the chemical guarantee of a sample of commercial oxide is expressed in the four ways in Table 11.

TABLE 11

**DIFFERENT METHODS OF EXPRESSING THE CHEMICAL GUARANTEE OF COMMERCIAL OXIDE**

| Material | Oxide Content,<br>per cent | Elemental<br>Ca or Mg,<br>per cent | CaO<br>Equivalent | Neutralizing<br>Power |
|----------|----------------------------|------------------------------------|-------------------|-----------------------|
| CaO      | 75                         | 53.3                               | 103               | 183.3                 |
| MgO      | 20                         | 12.0                               |                   |                       |

The figures representing the oxide content are the forms in which chemists usually express the quantities of calcium or magnesium in any material. In other words, in this particular sample the calcium in all its forms, expressed as CaO, was equivalent to 75 per cent of the material. The magnesium compounds, whether they were in the oxide, hydroxide, or carbonate forms, expressed as MgO, made up 20 per cent of the material by weight. In arriving at these values, the chemist makes a determination for total calcium and magnesium (53.3 per cent and 12.0 per cent, respectively, for this sample) and expresses the results in terms of CaO and MgO. The calcium oxide equivalent is nothing more than a term expressing both the CaO and MgO in terms of CaO. Since 1 pound of MgO is equivalent to approximately 1.4 pounds of CaO, the CaO equivalent of the MgO in the sample is 28 or  $20 \times 1.4 = 28$ . The total CaO equivalent becomes 103 or  $75 + 28 = 103$ . As already explained (p. 144), the neutralizing power is obtained by expressing the strength of the lime in terms of its CaCO<sub>3</sub> equivalent; in this case it is 183.3 ( $103 \times 1.78 = 183.3$ ).

In most states having chemical lime-control laws, it is customary to state the guarantee of commercial hydrates and oxides in terms of oxide content. This shows the amount of magnesium as well as cal-

cium present, which is an important consideration particularly where magnesium may be needed in the soil. In addition, the guarantee may carry one or more of the other forms of statement. The statement of the chemical analysis in terms of CaO equivalent or in terms of neutralizing power is convenient in that it is made in one figure and thus renders comparisons between different materials quite simple.

The method of stating guarantees for ground limestone is somewhat different from that for commercial oxides and hydrates. The quantities of calcium and magnesium carbonates are usually expressed separately and in terms of total carbonates on a percentage basis. These may also be expressed in terms of CaO equivalent, in terms of neutralizing power, or even in terms of elemental calcium and magnesium. The chemical guarantee of a representative sample of ground limestone may be expressed in one or more ways, as illustrated in Table 12.

TABLE 12

VARIOUS METHODS USED IN EXPRESSING THE CHEMICAL GUARANTEE  
OF COMMERCIAL GROUND LIMESTONE

| Lime              | Separate Carbonates | Carbonates, per cent | Neutralizing Power | Total CaO Equivalent | Ca or Mg, per cent |
|-------------------|---------------------|----------------------|--------------------|----------------------|--------------------|
| Ground lime-stone | 80% $\text{CaCO}_3$ | 95                   | 97.9               | 54.76                | 32.00              |
|                   | 15% $\text{MgCO}_3$ |                      |                    |                      | 4.32               |

Of the various methods of expressing guarantees, the neutralizing power (from which the CaO equivalent easily can be calculated) is the easiest and most rapidly determined. From the analytical point of view, this is an important consideration when many samples are involved. Yet the meaning of the carbonate method of guarantee is perhaps easier for the farmer to understand, and it gives the quantity of magnesium present—an advantage of this method of presentation. However, the differences between these two values (carbonates and neutralizing power) are usually so small that practically they are of little or no significance.

**Fineness Guarantee.** A fineness guarantee is based on a mechanical analysis of a particular sample of ground limestone made by the use of standard screens of definite mesh or size openings. A 10-mesh screen, for example, has 10 openings per linear inch or 100 openings per square inch. The quantities of stone that will pass through each of the various screens are stated in the guarantee. The following is an example: 100 per cent through an 8-mesh screen, 50 per cent

through a 60-mesh screen, and 40 per cent through a 100-mesh screen. Various grades of ground limestone appear on the market. They are based on the percentage of the stone that will pass through screens having a specified number of meshes per linear inch.

With all other factors equal, the finer a limestone is ground, the more rapidly it will dissolve and the more thoroughly it can be mixed with the soil. The effectiveness of a ground limestone of a given neutralizing power is determined not only by its rate of solubility but also by its contact with the colloidal particles. However, the finer the stone is ground, the greater will be its cost and the less will be its lasting qualities. Furthermore, a very finely ground limestone is not only difficult to handle but also unpleasant to distribute. Therefore, it is generally recommended that a ground limestone of medium fineness be purchased. Such a grade can be ground rather cheaply, and it contains a sufficient quantity of fine material to give immediate effects and a sufficient amount of coarse material to give it lasting qualities. Such a ground limestone would be one in which all would pass an 8-mesh screen and 25 to 50 per cent would pass a 100-mesh screen.

#### SOURCES OF LIME

In the introduction to this chapter, it was pointed out that one reason why calcium compounds rather than other compounds are used for liming purposes is that the supply of calcium compounds is plentiful and widely distributed. In this connection mention should be made of the different sources of lime and of the nature and supply of such materials. Questions are proposed to call attention to pertinent points relative to supplies of liming materials.

#### Questions

1. What is limestone? Is there a sufficient quantity to meet the needs of the soil?
2. What is marl? How is it formed, and where are the deposits found?
3. Is sugar-factory lime of any value for liming purposes?
4. Do wood ashes contain significant amounts of calcium?
5. Can lime from gasworks, acetone plants, acetylene plants, and paper mills be used for liming purposes?
6. Is blast-furnace slag of any value for correcting soil acidity?
7. Are oyster shells considered a satisfactory source of lime?

**Limestone.** Limestone is a sedimentary (deposited in water) rock formed by the precipitation of calcium and magnesium in water in

the form of carbonates with subsequent consolidation of the deposits. Limestone deposits are widely distributed, and they constitute an important source of lime. Many limestone quarries are operating in the humid regions of the United States and are in a position to meet the demand of agriculture, as well as of all other industries, for lime. It might be added that only a small portion of the output of these quarries is used in agriculture at the present time.

Limestone is the most important single source of commercial liming materials. The approximate consumption of liming materials on United States farms during 1948 is presented in Table 13. A very large

TABLE 13

## APPROXIMATE AMOUNTS OF LIMING MATERIALS USED ON UNITED STATES FARMS DURING 1948 \*

| Form of Lime                               | Tons Used  |
|--|------------|
| Agricultural limestone †                   | 23,763,903 |
| Hydrated lime                              | 174,034    |
| Marl                                       | 515,785    |
| Burned lime                                | 124,551    |
| Miscellaneous materials ‡                  | 232,937    |
| <hr/>                                      |            |
| Total materials                            | 24,811,210 |
| Lime oxide equivalent                      | 12,369,244 |
| Pounds of lime oxide per acre of crop land | 80.3       |

\* From data compiled by the National Lime Association.

† Includes all grades of limestone, varying from that which will all pass a 20-mesh screen to the coarser material, often called "screening."

‡ Includes egg and mollusk shells, paper- and sugar-mill refuse, lead-mine chats, and carbide refuse.

percentage of the commercial lime that is used for liming purposes is used in the form of ground limestone or in some form made directly or indirectly from limestone. The calcium of commercial hydrate, commercial oxide, and calcium silicate of the blast furnaces all comes from limestone.

**Marl.** Marl<sup>1</sup> or bog lime is formed by lime which was dissolved from the soil in drainage water and carried to swamps, lakes, and

<sup>1</sup> From an agricultural point of view bog lime is usually spoken of as marl. Geologists frequently refer to marl as a calcareous clay of variable composition. In certain areas unconsolidated calcareous rock is called marl; the calcium carbonate is finely divided and is also in the form of shells and fragments of shells. Glauconite (greensand) is also sometimes referred to as marl.

other bodies of water where it was thrown out of solution. This precipitation of calcium may be a purely chemical reaction, the loss of carbon dioxide from calcium bicarbonate, or it may occur as the result of the action of certain plants or of shell-forming animals. Frequently it accumulates on the bottom of these bodies of water as a soft, mush-like material, or it may occur in nodular form. Often these two forms are intermixed. It may be found underlying areas of muck and peat

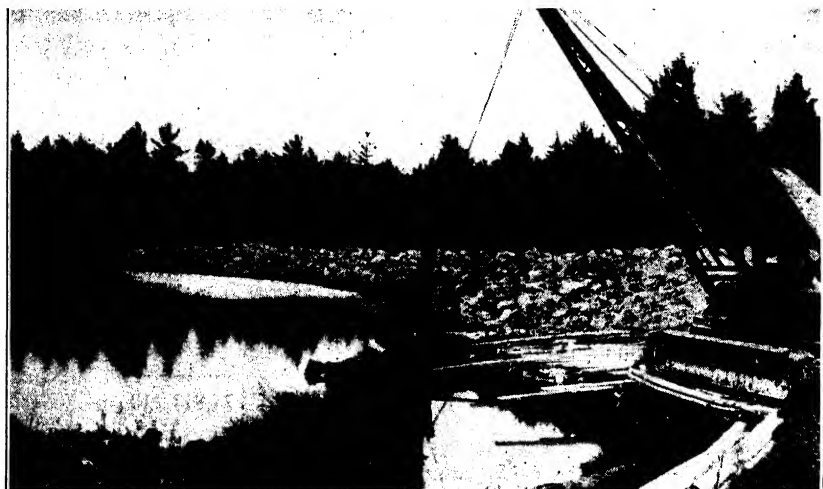


FIG. 30. A marl pit in Michigan.

and along the shores and in the beds of lakes. The marl may be covered by a few inches or several feet of muck. Marl beds are also found on dry land where an old lake has been drained by a change in the drainage level. The thickness of these marl deposits may range from less than one inch to several feet, and the areas are quite variable in size.

The lime in marl exists principally in the calcium carbonate form with varying quantities of magnesium carbonate. The value of a marl to correct acidity is usually expressed as though all the carbonates were present as calcium carbonate; thus a marl testing 95 per cent has the same neutralizing power as a ground limestone with a neutralizing value of 95. The purity of marl may vary considerably, even within the same deposit. The common impurities found in this material are sand, silt, clay, and organic matter. Of these, clay is perhaps the most objectionable. A marl containing a medium to low clay content is very

sticky when wet and becomes very hard and lumpy when dry; furthermore, it is difficult to handle and cannot be applied uniformly.

Owing to the variable moisture content of marl, it is usually spoken of and dealt with in terms of cubic yards. On the average, a cubic yard of good marl (testing about 95 per cent on the dry basis) weighs about 2,300 pounds in the wet condition and contains approximately 1,600 pounds of calcium carbonate. Since it is usually difficult to spread marl uniformly, 2 cubic yards of a good marl frequently are recommended for each ton of limestone needed.

In some states, marl is an important source of liming materials. In Michigan, for example, approximately 27 per cent of all liming materials used is in the form of marl.

**Sugar-Factory Lime.** In the manufacture of sugar from sugar beets, lime water is added to clarify and purify the juice from the beets. Carbon dioxide gas is then bubbled through the juice to precipitate the lime as calcium carbonate, which is subsequently filtered out and discarded. This material contains insignificant quantities of organic matter and plant nutrients. This lime frequently is washed out to the waste pile with the water used in washing the beets and may thus contain considerable dirt. On a dry basis, sugar-factory lime will usually test about 80 per cent  $\text{CaCO}_3$ . Owing to its variable moisture content, it usually is handled on the cubic-yard basis.

Some of the sugar factories prefer to burn their own lime, and, when the kilns are cleaned, impure burned lime or stone lime is obtained. This material is mostly  $\text{CaO}$  when fresh, but after a short time it takes on water and forms hydrate, and in turn some of it may react with carbon dioxide, forming calcium carbonate.

**Wood Ashes.** In certain localities wood ashes may be available in sufficient quantities for use on the soil. The composition will depend not only on the composition of the original wood but also on the completeness of burning, leaching that has occurred, and impurities with which it may have become mixed. The lime in freshly produced ashes is in the oxide form, but on exposure hydroxide and carbonate will be formed. Wood ashes, as ordinarily used, may be expected to contain from 20 to 50 per cent of lime, expressed as calcium carbonate. They will also contain about 5 per cent or less of potash and very small amounts of phosphorus but are usually regarded as more important as a liming material than as a source of potash.

**Lime from Acetone Plants, Gasworks, Acetylene Plants, and Paper Mills.** There are various manufacturing plants which produce or discard lime as a by-product. Such material is usually of rather high

grade, and, when it is available locally and at a low cost, its use may be advisable. The principal disadvantage of this lime is that it usually contains a high content of water, which makes spreading difficult. The cost of transportation and the difficulty in distributing must be taken into consideration when the costs are compared with those of other sources of lime. These materials generally can be obtained at little or no cost since they are regarded as waste products.

Calcium oxide is used at gasworks to remove impurities from the gas, after which it is either discarded or sold to farmers. This lime is largely in the hydroxide and carbonate form, and it contains many impurities, some of which, such as sulfites and sulfides, may injure seeds. On exposure to the air, these sulfur compounds soon change over to the sulfate form, which is harmless.

Acetylene lime is a solid residue formed in the manufacturing of acetylene gas from calcium carbide and consists largely of the hydroxide and carbonate forms of lime. This material is wet and contains some acetylene gas which is injurious to seeds, but this gas will escape within a few weeks on exposure to the air.

**Blast-Furnace Slag.** As mentioned previously (p. 144), blast-furnace slag contains calcium in the form of calcium silicate. The slag is a by-product in the smelting of iron ore. This may be considered a liming material inasmuch as it supplies active calcium, leaves no objectionable residue (the silicate ion produces a very weak acid), and has the ability to reduce soil acidity. Basic slags (in which materials are added to remove acid impurities) usually contain magnesium oxide and also calcium oxide. A typical grade of basic slag produced in the United States contains 6 to 8 per cent of calcium silicate, 5 to 7 per cent of magnesium oxide, and 45 to 48 per cent of calcium oxide. This is an important source of lime in some of the Southeastern States. Basic slag is also an important carrier of phosphorus (see p. 336).

A relatively new type of slag is now becoming available in substantial quantities from rock phosphate reduction furnaces. This product consists mainly of calcium silicate and contains most of the original constituents of phosphate rock, except the phosphorus, together with the silica which was added. This product constitutes a satisfactory liming material.

**Oyster Shells.** Clean oyster shells contain about 90 to 95 per cent of calcium carbonate. When finely ground, they are a very satisfactory liming material which is used to some extent on farms located near the source of supply. Oyster-shell lime sometimes is burned and used in the calcium oxide form.

### THE USE OF LIME

After a discussion of the general importance of lime in agriculture, its functions in the soil and the forms, composition, and sources of lime, it is important that certain points dealing more specifically with the practical use of lime be clearly understood. May it be emphasized again that lime, if needed and wisely used, is indispensable to a permanent agriculture. Several questions are suggested as guides in this discussion.

#### Questions

1. How can the need of soils for lime be determined?
2. Are all crops benefited by the application of lime?
3. What kind of lime should be used?
4. What changes does lime undergo when applied to soil?
5. How much, and how frequently, should lime be applied?
6. When should lime be applied?
7. What are the different methods of applying lime?

**Determining the Need of Soils for Lime.** In arriving at a conclusion about the desirability of applying lime and the amount needed, it is necessary that soil tests be made. The general need of soils for lime may be made by observation within certain limits, but the amounts to apply are not so indicated. Some common indications of an acid soil are: (1) a prevalence of red sorrel; (2) the repeated failure of alfalfa and clover; (3) thin or patchy stands of alfalfa, or clover on soils which produce good yields of other crops; or (4) alfalfa and clover may turn yellow in early spring. These symptoms may be caused by some factor other than a lack of lime; yet these conditions frequently are associated with a deficiency of calcium. From such observations it is obvious that the amount of lime needed cannot be ascertained, and it may be practicable to use it long before any of the deficiency symptoms are apparent. Furthermore, an excess of lime is to be avoided; consequently soil tests must be used. The idea should be discarded that lime is a cure-all and that, if a little is good, more is better.

Soil tests for determining lime requirement are discussed in Chapter 6 (p. 124) and will not be repeated here. It may be mentioned, however, that the simple colorimetric tests are sufficiently accurate for practical purposes. It is much more desirable from the practical standpoint to test a large number of samples from a particular field with the simple tests than to test only one or two samples by one of the more accurate but time-consuming laboratory methods.

The different fields on any particular farm frequently vary a great deal in acidity; likewise, each field may vary in acidity from place to place. Consequently, when the whole farm or an entire field is given a uniform application of lime, some parts may get too little and others too much. It is important, therefore, that a field be properly sampled and tested in arriving at the lime needs. Several tests should be made of the surface soil (plow depth), and at least one or two tests of the subsoil (at a depth of 2 to 3 feet) of each soil type in each field. It is sometimes recommended that the surface soil of each acre of a field be tested. The number of tests which should be made in a field of a given size will depend principally on the topography. On very level fields fewer tests need be made. It is necessary that several tests be made of each soil type in order to avoid such irregularities as may be due to droppings of manure, ashes from burned weeds, brush, or crop residue, and straw-stack residues.

In several states soil-testing laboratories have been established at various locations. Some of these serve one county, and others a larger area. The laboratories are sponsored by one or more of several agencies, as the Farm Bureau, the Board of Supervisors, directors of a Soil Conservation District, or other farmer organizations. The county agricultural agent, or farm advisor, frequently has direction of the laboratory, and results of tests are referred to him to be relayed and discussed with the farm owner or operator. Tests are made in these laboratories at a minimum charge. The soils or agronomy department of most state colleges or universities also maintain a soil-testing service. Most of the laboratories determine the lime needs of soils by means of portable glass-electrode sets.

Soil-acidity tests, together with an understanding of the chemical factors associated with the test results and an understanding of a plant's requirements, furnish a basis for making lime recommendations.

**Response of Crops to Lime.** Crop response to lime depends on a number of factors, among which the *pH* of the soil is an important one. It is, however, generally accepted that *pH* as such is rarely the limiting factor in plant growth within the range normally existing in agricultural soils. Plant growth frequently is retarded in acid soils because of unfavorable chemical conditions associated with a low *pH*; for example, the degree of base saturation is definitely related to *pH*, and, since calcium, magnesium, and potassium are all involved, plant growth must be affected by the supply of these elements. The solubility of aluminum, iron, manganese, and perhaps other minor elements is also related to *pH*, as is also the activity of microorganisms. The

experimental evidence relative to crop response to lime is, as would be expected, rather confusing because of the wide range of soil and climatic conditions under which many of the lime experiments have been conducted.

In general, the legumes respond more to lime than do the non-legumes. The fixation of nitrogen by legumes is much more effective



FIG. 31. Alsike clover is more tolerant of soil acidity than are alfalfa, sweet clover, and red clover. Nevertheless, on this field at the Tennessee experiment station the crop was greatly increased by an application of lime. [From U.S.D.A. *Farmers' Bull.* 1845.]

where high levels of available calcium are present. Of the legumes commonly grown, alfalfa, sweet clover, red clover, ladino, and Dutch clover respond to liming most markedly. Garden and field peas, soybeans, and alsike clover give noticeable response to lime, but less than the legumes previously mentioned. Likewise, cowpeas, vetch, field beans, and lespedeza are benefited to some degree by lime, but less than the preceding group. Contrary to popular belief, all legumes do not respond favorably to lime. For example, serratella and the lupines may actually be injured by liming.

Of the non-leguminous crops, sugar beets, table beets, cabbage, and cauliflower are particularly responsive to lime even though the soil may not be strongly acid. Corn, oats, wheat, barley, rye, sorghum, Kentucky bluegrass, and timothy appear to be benefited by liming, al-

though the benefit may be brought about in part indirectly. The indirect effect is produced through the stimulation of legume crops in the rotations which in turn benefit the non-legume crops. Lime permits greater growth of the legumes and thus leaves more organic matter and

larger quantities of active nitrogen in the soil. Crops like cotton, tobacco, Irish potatoes, redtop, strawberries, buckwheat, bent grass, fescue, and millet will make satisfactory growth on soils low in active calcium and appear to be little influenced directly by the addition of lime. However, indirectly (through its effect on legumes in the rotation) lime will usually have a beneficial effect. Certain non-leguminous plants may be injured by lime; included in the list are such plants as watermelons, blueberries, cranberries, laurel, azaleas, and rhododendrons. Furthermore, lime may injure potatoes by encouraging a disease known as scab.



FIG. 32. Although soybeans are somewhat tolerant of soil acidity, the crop responds remarkably to an application of lime on strongly acid soil. Left: Plants from limed soil. Right: The same number of plants from unlimed soil. [From U.S.D.A. *Farmers' Bull.* 1845.]

ing is a complicated phenomenon, and only rather broad, general conclusions can be drawn.

**Kind of Lime to Use.** In selecting the liming material to use, several factors must be taken into consideration which will vary with the conditions attending each individual farmer. The following factors are among those which should be considered in choosing the form or grade of lime that should be used: (1) relative cost of calcium in the

It is thus apparent that it becomes a matter of great importance not only to know the condition of the soil with respect to its content of active calcium and magnesium but also to understand the response of crops to these elements. As has been indicated, crop response to lim-

different forms, (2) fineness, (3) rapidity of action, (4) convenience in handling and storage, (5) character of soil, and (6) crops to be grown.

Since equivalent quantities of the different chemical forms of lime are essentially equally effective in neutralizing soil acidity and in increasing crop growth, obviously *cost* is one of the most important factors to consider in choosing a liming material. The cost of equivalent quantities of limes applied to the soil may be determined by simple arithmetic calculations. From these calculations it is easily learned which gives the greatest quantity of lime per dollar spent. Total cost is determined by the cost of the material at the plant or at the source of supply and the distribution costs, including delivery to the farm and application to the soil. If the material has to be transported a considerable distance, it may be less expensive to purchase the more concentrated materials (burned or hydrated lime). If the material is close at hand, the forms containing less calcium (ground limestone, marl, refuse lime, etc.) may be the least expensive.

Where immediate or first-year effects are especially desired, particular attention should be given to the degree of fineness, and under such circumstances allowance must be made for this factor in comparing costs of ground limestone, shells, etc. Properly slaked lime is probably in the finest state of subdivision of any of the liming materials. Hydrated lime is somewhat more soluble than the carbonate form, and until such time as it may change to the carbonate form the hydrate form must be considered more active.

The carbonate forms of lime are generally more convenient to handle and less disagreeable than the caustic forms, even when the latter are bagged. Furthermore, limestone in storage, if kept dry, will not change form as will the caustic forms. The difficulties in scattering materials, such as marl and certain industrial by-product limes, have been mentioned (pp. 153-154) and should be considered in selecting the form of lime to use.

Only in rather special cases does the character of the soil affect the form of lime that should be used. Obviously, on certain soils containing large quantities of magnesium, it is desirable to avoid the use of those forms of lime rich in magnesium. Furthermore, where the caustic effects of oxide and hydrate forms are likely to occur, such as on light, sandy soils, it is safer to use the carbonate form.

**Changes Lime Undergoes When Applied to the Soil.** The immediate changes that lime undergoes when added to the soil depend among other things on the form of lime, the chemical nature of the soil,

and the moisture supply. Lime in the form of either the oxide or hydroxide undergoes rather rapid change when applied to moist soils. The attraction between burned lime (oxide form) and water is very intense. Water combines with the oxides of calcium and magnesium to form the corresponding hydroxides or water-slaked lime. In moist soil this reaction occurs so rapidly that perhaps the oxides as such do not have time to react chemically with the soil.

Slaked lime has a strong affinity for carbon dioxide and the two combine to form carbonate. In moist soil the reaction probably occurs as follows:



This reaction would proceed rapidly in moist soils well supplied with active organic matter because of the abundant production of carbon dioxide.

It is thus evident that lime applied to soils in the form of oxide or hydroxide is sooner or later changed to the carbonate form. It might thus be assumed that little difference would be found in the activity of the various forms since they all change to the carbonate form. This is essentially true; yet, as has been mentioned before, the slaked limes are more soluble than the carbonate forms, and until they are converted to the carbonate forms they will be more active. Furthermore, it should be remembered that lime carbonate produced by the reaction of hydrated lime and carbon dioxide is in a physical state which dissolves more easily than do the dense particles of ground limestone, even though they be very small.

However, under field conditions over a period of years differences in the effectiveness of the various forms of lime generally have not been observed and practically are of little significance. Lime (in its various forms) may be considered to act principally in the carbonate form in the soil.

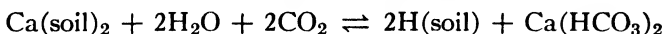
The carbonate form is only slightly soluble in water; yet in the finely divided state it appears to displace hydrogen ions of the colloidal soil particles, thus reducing the potential acidity. The reactions of the carbonate in neutralizing acidity may be illustrated as follows:



First the carbonate may react with carbonic acid to form calcium bicarbonate,  $\text{CaCO}_3 + \text{H}_2\text{CO}_3 \rightarrow \text{Ca}(\text{HCO}_3)_2$ , which in turn may react with the soil, or else the calcium carbonate may react directly with the

soil colloids. The colloids are thus charged with bases accompanied by the liberation of carbon dioxide.

It is obvious, therefore, that lime added to soils exists principally in two forms, as carbonates and as exchangeable ions in association with the organic and inorganic colloids. In both forms the bases are susceptible to the action of carbonic acid and form bicarbonates which may readily leach out of humid region soils. The reactions involved are essentially the reverse of those given above.



**Quantity and Frequency of Lime Application.** The amount of lime to be used depends primarily on the (1) degree of acidity of the soil, (2) buffer capacity of the soil, (3) acidity of subsoil, (4) crops to be grown, (5) form or grade of lime to be used, and (6) frequency of application. Various tests (p. 124) are used to aid in determining the amount of lime to apply to soils, but, because the quantity needed is dependent on so many factors, all lime recommendations are at best only approximations.

Under most general farming conditions, it is usually advisable to regulate the application of lime in order to maintain the soil reaction at a *pH* of about 6.5 or 7.0. Under other farming conditions and in special cases, it is sometimes desirable to regulate the quantity of lime so as to maintain either a higher or lower *pH*.

Sandy soils usually require less lime to bring them to any desired *pH* than do heavy soils having the same degree of acidity. This difference is due largely to the much greater buffer capacity of the heavy soils. Furthermore, the amount of lime required to change the *pH* of a strongly acid muck or peat soil is much greater than for mineral soils. On mineral soils it is usually not necessary and frequently impractical to apply more than the equivalent of 2 or 3 tons of finely ground limestone per acre or 3 to 4 tons of a coarser grade. Five to 10 tons of lime per acre are sometimes recommended for certain acid organic soils. Perhaps the most generally recommended application of lime considered necessary for mineral soils is 1,000 to 2,000 pounds of finely ground limestone (or its equivalent) per acre, once in a rotation. There is some difference of opinion about whether a large application is better than smaller and more frequent applications. When finances available for liming are limited, the latter is generally advisable.

In some instances, the use of a relatively small amount (200 to 500 pounds per acre) of very fine lime, drilled in the row at seeding time, has proved satisfactory. When this method is used, it is done as a regular part of the legume seeding whenever the crop comes around in the rotation. This method has an advantage in decreasing the initial cash outlay, gives a better distribution of labor, and permits the lime to be transported greater distances. It is not to be inferred that this method should displace the ordinary plan of liming, but the use of relatively small amounts of fine limestone may permit liming under many conditions where heavy liming is not commonly practiced.

The question of overliming has recently received considerable attention. It is generally believed that, if the *pH* of the soil is raised above 7.0 or 7.5, the availability of phosphorus, potassium, iron, manganese, and perhaps copper, boron, and zinc is decreased. There is evidence to indicate that in sandy soil with a decidedly low *pH* value, when the quantities of manganese, zinc, iron, and potassium are small, even moderate liming may result in a temporary deficiency of one or more of these nutrients. There is also evidence to indicate that overliming disturbs temporarily the utilization of phosphate in plants. No doubt, the question of overliming has in some instances been overemphasized, yet it is well to keep in mind some of the possible dangers resulting from such a practice.

**Time of Applying Lime.** Generally speaking, lime may be applied any time during the year when it is most convenient. Often, however, the type of the rotation, system of farming, and form of lime used will be the deciding factors.

If the caustic forms are to be used for spring-seeded crops, it is usually best to apply the lime in the fall. These forms will change over to the carbonate form by seeding time and will lose their caustic properties and have no injurious effect on seed germination. Lime in the carbonate form may be applied any time without danger of injury. Regardless of the form of lime used, it is best applied when the soil is not too wet; otherwise it is difficult to get an even distribution. Under wet-soil conditions, the caustic forms and very finely ground carbonate tend to ball badly and not mix evenly through the soil.

It is advisable, of course, to apply lime where it can be used to the best advantage in the rotation, for example, preceding the legume crop or in connection with a green-manure crop. It is usually best to apply lime a considerable time in advance of seeding legumes, in order that the lime will have time to correct the acid condition of the soil. To in-

sure best results, lime should be in the soil at least six months prior to seeding legumes, although successful legume seedings sometimes are obtained by applying lime immediately preceding or with the legume seeding.

If the legume is to be seeded with a nurse crop, such as wheat, frequently the lime is applied just before the wheat is seeded in the fall. In a rotation of corn, oats, wheat, and clover, it is usually most conveniently and satisfactorily applied before the wheat. If a sod crop is to be plowed under for such crops as corn or beets, to be followed by one or two grain crops in either of which a legume is to be seeded, lime probably can be most conveniently applied to the sod before plowing.

When potatoes are included in the rotation, lime should follow this crop if scab is prevalent, for it is generally believed that lime makes conditions more favorable for the development of the organisms which produce scab. Many potato growers have found that this crop gives greater yields when it follows a leguminous sod. Generally the increase in yields is sufficient to justify using lime on sour soils to make the growth of leguminous crops possible. This, therefore, usually necessitates the use of a minimum amount of lime directly after the potato crop is harvested.

In general practice it is usually recommended that lime be applied when most convenient since the most important consideration is that of applying it regularly in the rotation as needed. Late summer or fall application is usually most convenient. At this time of the year, there is a lull in farm work, and the roads and fields are in a good condition for hauling and spreading.

**Methods for Applying Lime.** The principal requirement of any method of applying lime is that it should be distributed evenly and, except when applied to pastures, it should be thoroughly mixed with the soil. Lime even as it dissolves moves to no appreciable extent horizontally and only to a limited extent vertically. Movement is not sufficient to distribute the lime evenly over the field or to mix it thoroughly with the soil. Since soil acidity is due largely to the colloidal clay acids, it is essential that lime come in contact with all the soil particles so far as possible. This requires a thorough and even mixing of the lime with the soil. An even distribution may be accomplished by the use of most of the standard lime spreaders on the market. The only way of mixing the lime thoroughly with the soil is by tillage operations.

There are three types of spreaders in common use: (1) trucks with specially designed hoppers and distributors, (2) the two-wheeled box type of spreader which can be operated by one man and a team, and (3) the endgate spreader, which is attached to the rear end of the wagon box. The distributing device of this attachment is operated by a chain that is running on a gear bolted to one of the rear wagon



FIG. 33. The two-wheeled box type of lime spreader is an efficient implement for applying ground limestone and hydrated lime. [From U.S.D.A. *Farmers' Bull.* 1845.]

wheels. It requires two men to operate the endgate spreader, one to drive while the other shovels the lime into the spreader. Similar types of attachments for manure spreaders are also on the market.

Grain drills sometimes are used for applying lime. The lime is put into the fertilizer compartment and may be drilled either at the time of seeding or as a separate operation. This is an excellent method of applying small amounts of finely ground limestone.

The old method of spreading lime from a wagon with a shovel has the advantage of not requiring any additional investment in equipment. Sometimes the lime is placed in piles over the field and later scattered with a shovel. These methods require more labor, are usually more expensive, and are more disagreeable; also, it is more difficult to

spread the lime evenly than to apply with a machine, but such methods can be used if a mechanical spreader cannot be obtained.

Marl, sugar-factory lime, and similar types of liming materials which

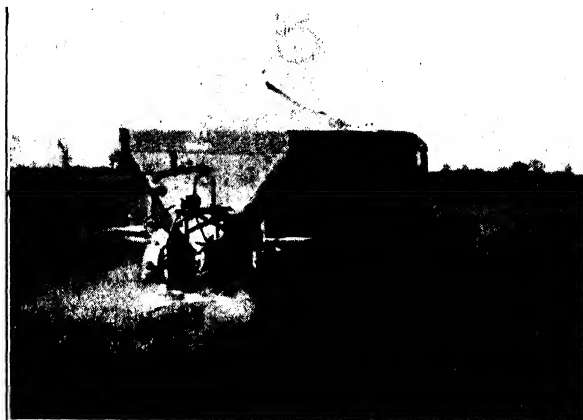


FIG. 34. Special trucks, with V-shaped beds having a screw conveyor in the bottom that carries the lime to a centrifugal spreader at the rear, are in common use.

are usually wet and difficult to handle are generally applied by means of a manure spreader or truck equipped with a spreading device.

A thorough mixing with the soil can perhaps be best accomplished by applying the lime evenly on plowed land. The disking, harrowing, and other tillage operations permit a thorough mixing of lime and soil particles.

# 8

## Soil Moisture

Moisture supply receives more general attention than any other factor concerned with crop production. The bringing of water to otherwise fertile land has entailed the construction of immense dams and elaborate canal and pumping systems and has resulted in the conversion of huge expanses of desert into highly productive farms. The use of irrigation to supplement natural moisture supply in humid regions also has been developed to a considerable extent, principally for the production of special crops. On the other hand, in humid regions the moisture problem is more often one of removal than of supply. In many places extensive drainage systems have been necessary to remove the excess water from countless acres of farm land in regions of moderate to high precipitation.

An inadequate moisture supply frequently limits crop production to a greater or lesser extent in virtually all regions where natural precipitation is depended on to supply the moisture needs of plants. The problem of getting sufficient water to enter the soil confronts many farmers. On irrigated land the farmer is interested primarily in increasing the permeability of his soil to irrigation water, and the farmer in humid areas endeavors to increase the percentage of the precipitation that enters the soil instead of running off over the surface. The farmers of both regions are concerned with the conservation of soil water for crop use.

As the moisture problem under irrigation will be discussed in another chapter, let us now consider moisture relations in humid and semi-humid regions. Water which reaches the earth in the form of precipitation may be classified under headings of (1) that which does not enter the soil and (2) that which does enter the soil. It is suggested that the study of soil moisture be taken up under five headings or objectives.

### Objectives

- A. Soil water which yields to the pull of gravity.
- B. Soil water which is retained against the pull of gravity.
- C. Water in relation to plant growth.
- D. Loss of moisture from the soil.
- E. Runoff water.

### SOIL WATER WHICH YIELDS TO THE PULL OF GRAVITY

Water which the soil is unable to retain against the force of gravity is known as gravitational, free, excess, or drainage water. As this moisture occupies the larger spaces in the soil which should be filled with air, if plants are to grow well, the rapidity with which gravitational water passes below the main root zone of crops is of interest. Soil through which free water passes slowly is referred to as tight or cold, or as having poor internal drainage. Planting frequently is delayed on such soils, with the resulting hazard of frost damage in the fall in some climates. In other soils the movement of free water is very rapid, producing warm or early soils. The escape of gravitational water usually is influenced by the underlying soil horizons to a greater extent than by the surface soil, and hence when soil drainage is under investigation an examination should be made to a depth of three or more feet. In moving downward through the subsoil, the drainage water as a whole follows well-developed drainage channels and does not seep slowly around particle after particle. There is, of course, some downward and oblique seepage as the moisture moves toward the channels of freer flow. Likewise, as the channels fill there is seepage out from them so that the entire body of the soil is wet. These drainage ways are not large and are comparatively close together. They consist of channels left by decaying roots and of those made by worms and insects. Small crevices between the structural units of the soil also serve as drainage channels, as do numerous cracks formed by drying of the soil. An idea of the methods used in removing excess water from the soil and of the benefits to crops of its removal may be gained by seeking answers to the questions proposed.

### Questions

1. Why do many crops grow poorly in inadequately drained soils?
2. How should tile lines be placed in the field?
3. How deep and how close together should tile lines be placed?

4. What size of tile is advisable?
5. Should careful attention be given to outlets?
6. Is tile drainage always preferable to open ditches?
7. Are appreciable quantities of nutrients lost in drainage water?

**Detimental Effects of Poor Drainage.** Poorly drained soils warm up slowly in the spring and cool slowly in the fall because water has a heat-absorbing capacity approximately five times greater than soil particles, and therefore much heat energy must be added or lost to bring about a change in temperature. The early-season coldness of poorly drained soils is one of the limiting factors in their use for crop production. The high moisture content of these soils also delays the fitting of the soil for planting and thus necessitates their use for late-planted crops or for the seeding of early crops in poorly prepared soil.

The roots of the most commonly grown crops require a supply of oxygen for the process of respiration, and as poorly drained soils contain an abnormally low proportion of air the roots tend to suffocate in them.

With an inadequate oxygen supply, decomposition of organic matter must be largely of an anaerobic nature with the production of compounds, some of which are toxic to plants, in a reduced chemical state. The disagreeable odors from stagnant pools during warm weather bear witness to the production of unoxidized compounds in waterlogged soil. Under such conditions oxidized mineral compounds may be also reduced to poisonous combinations such as sulfides from sulfates, nitrites from nitrates, ferrous iron from ferric iron. Plants may also suffer from an insufficient supply of nutrients in poorly drained soils because the normal decay of organic matter with the accompanying liberation of available nutrients is hampered. Furthermore, it is doubtful that the plant roots can effectively carry on the processes by which nutrients are liberated from the surfaces of the soil particles and are absorbed.

**Systems of Tile Drains.** Different soil situations require different types of drainage systems. A few of the conditions frequently encountered will be discussed. First, there are the level lands with a high water table which require a complete, systematically placed tile system. The main tile lines should be placed at such intervals that the secondary lines, commonly called laterals, flowing into them will not be more than 80 rods long. Furthermore, the laterals should enter the main lines at a small angle to the direction of flow rather than at right angles to it. Two commonly used systems for such land are shown in Fig. 35.

Rolling land of clay loam or silt loam soil may need drainage only in the valleys. The first inclination is to place the tile lines near the center of the valley, as that is where the need for drainage is most ap-

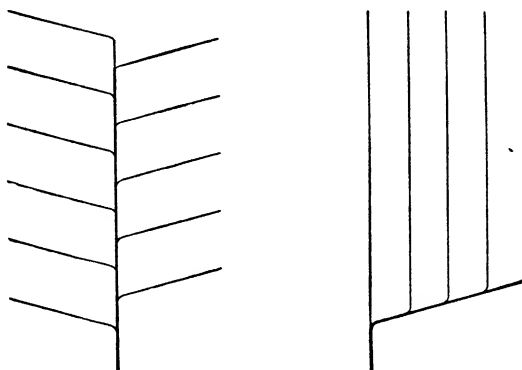


FIG. 35. Two systems used in laying tile. On the left is shown the herringbone method and on the right the gridiron system. The larger tile line into which the smaller ones or "laterals" empty is called a "main."

parent. It is usually found, however, that the water seeps down from the adjoining hills, and the proper place to install the tile is near the foot of the slope, where they will intercept the water and prevent it from reaching the main valley floor. The aid of an experienced drainage engineer is needed for the proper location of effective drains in such land. The depth at which such drains are laid is highly important.

A covering of sandy soil of varying depth over clay frequently offers an interesting drainage problem. The portion of the field where the sand is of considerable thickness may need no drainage, but as the sand covering gets thinner excess water is likely to cause trouble. Again, in such soil the tile should be placed not in the main body of the wet sand but near the edge of the thicker sand covering, for it will be found that the water is moving over the clay floor from the deep sand. The tile should also be set so that they lie largely in the clay.

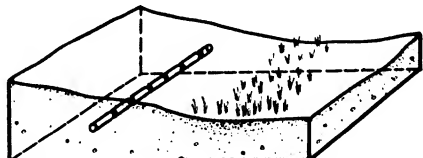


FIG. 36. To drain a wet or "springy" area near the bottom of a slope, the tile is not put through the poorly drained spot itself but some distance up the slope so that it will intercept the water as it seeps downward.

In some fields comparatively small depressions are wet although the soil as a whole is sufficiently well drained. It would seem that lines of rather small tile would be adequate to remove the excess moisture from such small areas. This conclusion is incorrect, since the water from a considerable portion of the field may slowly accumulate in these depressions. Accordingly, it is necessary to use a sufficiently large size of tile in each case to carry the water from all the land that drains into the depression.

**Depth and Spacing of Tile Lines.** Unless some unusual condition prevails, tile should be laid at least  $2\frac{1}{2}$  feet deep. The depth of the outlet has some bearing on the depth at which the drains may be placed and still provide a satisfactory fall. A fall of at least  $\frac{1}{10}$  foot per 100 feet is considered necessary, and more is desirable. The water level is lowered by the lines as shown in Fig. 37. In order, therefore,

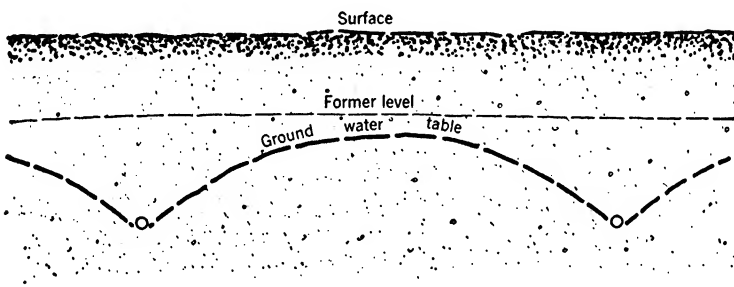


FIG. 37. The effect of tile lines in lowering the water table or groundwater level. In very heavy soil it takes several years for the water table to reach the proper adjustment. The benefit of the drainage is first evident immediately over the tile and gradually spreads to the soil area between them.

to have a sufficiently low water table between tile lines, to permit crops to develop an adequate root system, it is essential that the tile be  $2\frac{1}{2}$  or more feet deep.

The freedom with which water moves through the soil determines the distance apart drains may be placed and yet afford adequate drainage. If the soil through which the water must percolate to reach the tile contains a high percentage of clay and the structural condition is such as to make water movement slow, the tile lines should not be more than 4 rods apart. In more pervious soil a spacing of 5 rods is permissible, and in soil through which water drains rapidly the drains may be placed at 6-rod intervals. If the entire field needs drainage, tile lines should not be farther than 6 rods apart even in very pervious soils.

**Sizes and Kinds of Tile.** Experience and study have led to the conclusion that the small 2- and 3-inch tile formerly used extensively for laterals are not advisable. Nothing smaller than a 4-inch tile is now recommended even for short lines, and, if the drain is to be of considerable length, a 6-inch size is preferable. Main lines into which the laterals empty should be of 6-inch, 8-inch, or larger tile, depending on the volume of water to be carried. In the long run it is better to use tile a size larger than is thought necessary rather than a smaller size. A drainage engineer will have a table for calculating the size of tile to use according to volume of water to be carried, fall, rate of movement of water into the tile, and similar factors.

Three kinds of tile are usually found on the market. The soft clay tile are of varying quality. If well-burned so that they give a clear, ringing sound when struck, they are quite satisfactory. On the other hand, if they are poorly burned they are usually not very durable. Tile which are glazed, generally by the use of salt, are of very good quality and, although more expensive, are more economical than a poor grade of soft clay tile when length of service is considered. For a number of years concrete tile have received considerable attention. They vary greatly, owing to the difference in curing procedure. Unless the manufacturer has facilities for keeping the tile moist for a time sufficiently long to permit thorough setting of the cement, they are not durable. This uncertainty in quality has brought concrete tile more or less into disrepute. Also, in highly acid soils they are likely to disintegrate prematurely. They are seldom recommended for use in muck or peat soils.

In laying drains in organic soils or in other soils which are not sufficiently rigid to form a firm base and so keep the tile in line, it has been customary to place them on boards to prevent settling. A better procedure is to use a somewhat larger tile and procure them in double length (2 feet). Each of these tile will present much more bearing surface than the short ones of smaller diameter, and hence they will stay in line much better.

**Good Outlets Important.** Before a drainage system is installed, a careful investigation should be made of outlets which will provide for disposition of the water removed from the land. Frequently, when there are extensive areas of land which need drainage, the natural streams which can serve as outlets are so far apart that tile lines must be too long. They would be required to carry such a large volume of water that very large and expensive tile would be needed, and also it would be impossible to get sufficient fall and yet place the tile at the upper end of the system deep enough to provide efficient drainage. Also, many

such natural streams are too shallow to afford good outlets for tile lines and too small to carry the volume of water which would be quickly thrown into them from a tile system. To overcome these limitations dredging of the natural streams is frequently resorted to and, in addition, a quite complete system of dredge ditches is sometimes necessary.

In some land areas lying close to lakes, bays, or large streams into which the drainage flows, there is not sufficient difference between the land and water levels to allow for satisfactory tile drainage. Such situations require the most careful investigation by competent drainage engineers before steps are taken to develop the land. Some attempts have been made to dike areas of such land when the quality of the soil appeared to warrant so expensive an enterprise. A drainage system would be installed which carried the water to concentration points along the dike, from which it would be pumped over the dike into outlet streams or ditches. The expense involved in the construction and operation of such developments makes their feasibility questionable except under very unusual conditions.

The openings of tile lines into streams and ditches should be constructed in such a way as to prevent the undermining of the outlet by water and the slumping of the tile out of line. Several lengths of sewer tile at the end of the drain are often suggested. The opening of the tile should also be protected against the entrance of rabbits and other small animals.

**Tile Lines vs. Open Drains.** A tile drainage system has several advantages over drainage by open ditches, some of which are (1) when properly installed, little cleaning or other maintenance work is required; (2) the land over the tile is available for crop production, and the inconvenience of turning at the ditch when working in the field is avoided; (3) there is no ditch bank to serve as an area for production of weed seed; and (4) tile lines may be placed at closer intervals in the field and so provide more complete drainage. Two disadvantages of tile systems are (1) the initial cost and (2) their inability to carry an excess volume of water during periods of unusually heavy rainfall.

In fields which are inadequately drained with tile or open ditches, surface drains made by running dead furrows across the field at frequent intervals help to remove the excess water during heavy spring rains. These drains are usually closed by cultivation later in the season.

**Loss of Nutrients in Drainage.** As previously mentioned, drainage water tends to displace the soil solution from the surface soil and small cracks and channels as it passes downward through the soil. This phenomenon may be demonstrated by filling a percolation cylinder with

moist fertile soil and allowing it to stand for several hours to permit the soil solution to come to equilibrium with the soil. An inch or so of distilled water is now added to the surface and the percolate caught at the bottom of the tube in successive portions of 25 ml. Conductivity determinations on these portions will show that they contain appreciable and quite uniform quantities of soluble salts until the distilled water



FIG. 38. Exterior view of lysimeters in use at the Virginia Agricultural Experiment Station.

added to the surface approaches the bottom of the percolator. It becomes evident then that drainage is a source of loss of plant-food materials and would be entirely objectionable were it not for the fact that the space occupied by excess water in the soil is needed for air.

Table 14 presents results from the Connecticut Experiment Station. The lysimeters were filled with Merrimac sandy loam, a soil used extensively for tobacco growing in the Connecticut Valley. It is noted that growing a tobacco crop materially reduced the leaching loss of all nutrients except sodium. The application of the nitrogen fertilizer greatly increased the loss of sodium and nitrogen. An oat cover crop served to reduce drainage losses of all nutrients, although the saving of sulfur was small.

TABLE 14

## AVERAGE ANNUAL LOSSES OF NUTRIENTS IN DRAINAGE WATER

| Soil Treatment *                      | K               | Na   | Ca   | Mg   | S    | Cl   | N    |
|---------------------------------------|-----------------|------|------|------|------|------|------|
|                                       | Pounds per acre |      |      |      |      |      |      |
| No nitrogen (not cropped)             | 95.4            | 7.7  | 83.3 | 18.3 | 65.0 | 20.5 | 53.4 |
| No nitrogen (tobacco)                 | 75.0            | 7.5  | 57.6 | 15.1 | 57.8 | 15.3 | 27.1 |
| Nitrogen (tobacco)                    | 70.0            | 62.5 | 55.4 | 17.1 | 53.0 | 17.3 | 74.2 |
| Nitrogen (tobacco and oat cover crop) | 47.3            | 53.7 | 51.6 | 11.7 | 52.4 | 13.4 | 32.1 |

\* All lysimeters received a basic treatment per acre of 100 pounds of P<sub>2</sub>O<sub>5</sub>, 200 pounds of K<sub>2</sub>O, and 50 pounds of MgO in the forms of precipitated bone, carbonate of potash, sulfate of potash, and magnesium carbonate. The nitrogen treatment consisted of 120 pounds of N in cottonseed meal and 40 pounds of N in each castor pomace, and nitrate of soda.

The quantities of nutrients lost in drainage water are extremely variable. Efforts to determine the quantities of different nutrients lost from a particular soil under given climatic conditions and when a specific cropping sequence and soil treatment system are followed have been made by use of lysimeters. They are concrete or metal tanks of various sizes, set so that the top extends a few inches above the ground and having a bottom so constructed that the drainage water will flow through a tin tube into a receptacle. Two rows of tanks usually are set a suitable distance apart so that a passageway may be constructed between them to hold the drainage receptacles and allow space for the attendant to measure and sample the percolate. The inside of the lysimeters is coated with asphalt or other water-proofing paint which is not toxic to plants. The tanks are filled with soil taken from the field in 6-inch or 1-foot layers and placed in them in the order in which the layers occurred in the field. Some tamping is necessary to get the full depth of soil into the lysimeter.

The lysimeter method has been used to study the quantity and composition of drainage water by a number of experiment stations. Some of the larger installations are at the experiment stations in Illinois, Tennessee, New Jersey, and New York. The inside dimensions of the tanks at the New York Experiment Station, Ithaca, are 4 feet 2 inches square

by 4 feet deep. They are filled with  $3\frac{1}{2}$  tons of silty clay loam soil which was removed from the field and placed in them in 1-foot layers. Some of the lysimeters are cropped to different rotation systems and others are uncropped.

The effect of cropping on the loss by leaching is worthy of consideration, as are also the large quantities of certain ions lost, especially  $\text{Ca}^{++}$ ,

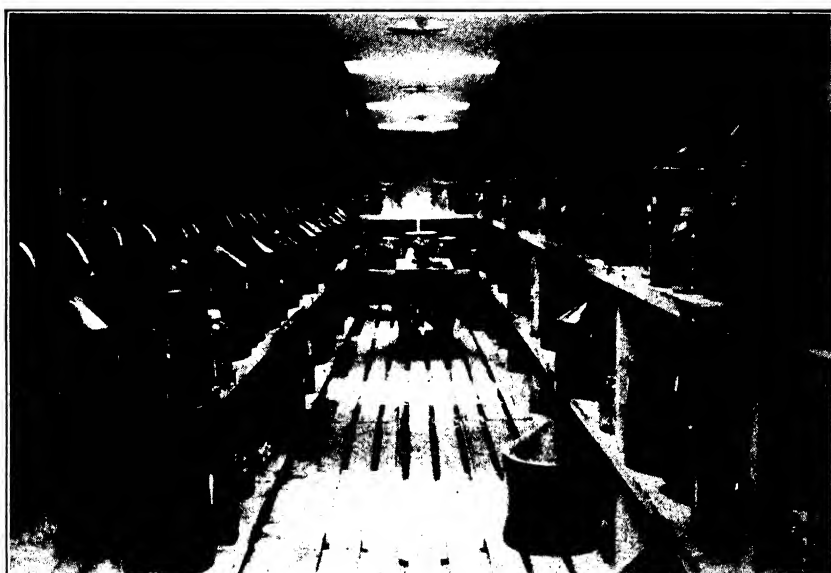


FIG. 39. Interior view of lysimeters at the Virginia station. Note receptacles for catching percolate which comes from lysimeter tanks through tin tubes.

and  $\text{NO}_3^-$ . Some of the factors which affect loss of nutrients in drainage water have been found to be (1) whether or not the soil is cropped; (2) the cropping system; (3) annual rainfall; (4) application of fertilizer, lime, or manure; (5) soil texture; and (6) soil composition.

The effect of cropping land, compared to keeping it cultivated but not planted to crops, on the total quantities of the various nutrients removed from the soil is illustrated by the data in Table 15, which are from the New York Experiment Station at Ithaca. It is seen from these results that the production and harvesting of a crop may actually result in the removal of appreciably less  $\text{Ca}$ ,  $\text{Mg}$ , and  $\text{NO}_3$  than if the land were clean-cultivated (fallowed) throughout the season. These results suggest that summer fallow should only be resorted to when necessary

TABLE 15  
EFFECT OF CROPPING ON THE REMOVAL OF NUTRIENTS FROM THE SOIL \*

|  | Potassium |           | Calcium |           | Magnesium |           | Sulfur  |           | Nitrate ( $\text{NO}_3$ ) |           |
|--|-----------|-----------|---------|-----------|-----------|-----------|---------|-----------|---------------------------|-----------|
|  | Cropped   | Uncropped | Cropped | Uncropped | Cropped   | Uncropped | Cropped | Uncropped | Cropped                   | Uncropped |
| Removed in the harvested crop                          | 78.8      | ...       | 13.2    | ...       | 7.1       | ...       | 10.1    | ...       | 77.0                      | ...       |
| Lost in drainage                                       | 45.9      | 61.0      | 173.3   | 367.4     | 37.0      | 65.2      | 38.2    | 48.5      | 5.4                       | 92.8      |
| Total  | 124.7     | 61.0      | 186.5   | 367.4     | 44.1      | 65.2      | 48.3    | 48.5      | 82.4                      | 92.8      |
| Reduction of loss in drainage as a result of cropping  | 15.1      | ...       | 194.1   | ...       | 28.2      | ...       | 10.3    | ...       | 87.4                      | ...       |
| Decrease in total loss to soil as a result of cropping | -63.7†    | ...       | 181.9   | ...       | 21.1      | ...       | 0.2     | ...       | 10.4                      | ...       |

\* Values expressed in pounds per acre per year.

† This means that 63.7 pounds more of potassium are removed from the cropped soil than from the uncropped soil.

for the storage of moisture for a following crop or for the eradication of noxious weeds.

### SOIL WATER WHICH IS RETAINED AGAINST THE PULL OF GRAVITY

If the pore space in a soil is filled with water and then drainage is permitted until no more water yields to the pull of gravity, the quantity remaining in the soil is known as the *water retained* or *field capacity*. Plants draw very largely on the retained water for their moisture supply, and hence the capacity of a soil to hold water against the pull of gravity becomes of practical significance. Not all the retained water is available to plants because the mutual attraction between soil and water is so great that a certain portion of moisture is held with a force greater than the absorptive power of plants. Several questions are suggested as guides in the study of this section.

#### Questions

1. What forces hold water in soil?
2. How can the forces holding soil moisture be measured?
3. How is the moisture content of soil expressed?
4. By what methods may soil moisture content be determined?
5. Into what classes may retained water be divided?
6. How does moisture move in the soil?
7. Does moisture move in the vapor state?

**Forces Holding Water in Soil.** Two forces are primarily responsible for the retention of water in the soil against the pull of gravity: (1) the mutual attraction between soil and water, which may be designated as adhesion; and (2) the attraction of water molecules for each other, which is a manifestation of cohesion.

If there were no attraction between soil and water, there would be no force to hold water in the soil and it would all yield to the pull of gravity and drain away. In other words, the retention of water in the soil is primarily due to the force of adhesion. With this retaining force in operation, the quantity of water held and the forms or shapes assumed by the water films are due largely to cohesion, as manifested by surface tension, which is a phenomenon resulting from unbalanced cohesive forces. Surface tension is illustrated in Fig. 40. In this illustration any molecule in the main body of the liquid, such as molecule *A*, is being attracted equally by molecules all around it, and hence its position is the result of balanced forces of cohesion. On the other hand, molecules

at the surface are attracted mainly by molecules to each side and below them. Consequently, the resultant force is downward, resulting in a compression and so a higher concentration of water molecules at the surface than in the body of the liquid. These molecules act as though they formed a tightly stretched elastic membrane over the surface of the water. This is the phenomenon known as *surface tension*.

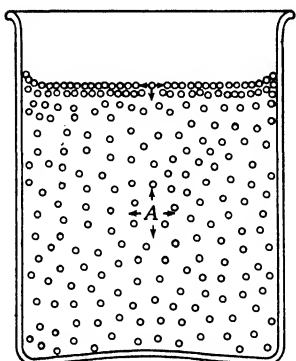


FIG. 40. A concentration of molecules at the surface of a liquid, as a result of unbalanced cohesive force, acts as a tightly stretched elastic membrane.

The apparent inclination of molecules to climb up the side of the beaker is evidence of adhesion and shows that the attractive force between glass and water is somewhat greater than the attraction between the water molecules themselves. A similar relationship exists between water and soil particles.

Temperature has a marked influence on cohesion in water and hence on the quantity of water retained. This fact may be demonstrated by permitting a column of saturated soil to drain until outflow ceases in a low temperature and then removing the column to a warm room and noting that drainage will again take place. The presence of soluble organic substances in the soil solution or of soluble salts will alter the surface tension of the solution and so affect the quantity of water retained.

#### **Measurement and Expression of Forces Holding Water in Soil.**

There has been some effort to express the force by which water is held in soils at different moisture contents in terms of centimeters in height of a column of water required to produce a force of equal magnitude. It has been proposed that the logarithm of the number of centimeters be used as an expression of the energy involved and that the value be called the  $pF$  of soil moisture. The  $p$  is to indicate that the value is a logarithm, and  $F$  is a symbol of free energy. The expression has some similarity to the designation of soil reaction in terms of  $pH$ .

The energy with which water is held in soils at different moisture contents may be measured by several methods. A very high-velocity centrifuge is made which will submit the soil moisture to pulling forces as high as 300,000 times gravity. Perhaps the most widely used apparatus is the tensiometer. This consists of a porous cup made of clay

and attached to a mercury manometer. The cup and connecting tube are filled with water, and the cup is placed in the moist soil. When equilibrium is reached between the moisture in the cup and the soil, the attractive force of the soil for water at that moisture content may be determined by reading the height of the mercury in the manometer. The values so determined are sometimes called the *capillary potentials* of the soil. Determination of the freezing point of a soil at different moisture contents is another way of determining capillary potentials. The capillary potentials of a sand, loam, and clay at various moisture contents are given in Table 16. The great variation in moisture content

TABLE 16

THE CAPILLARY POTENTIAL OF SOILS OF DIFFERENT TEXTURES AND AT DIFFERENT MOISTURE CONTENTS \*

| Soil | Moisture Content in Percentage of Dry Soil—Capillary Potential in Centimeters per Gram |      |      |      |      |      |      |      |      |      |      |
|------|--|------|------|------|------|------|------|------|------|------|------|
|      | Moisture content   | 20.9 | 11.7 | 7.6  | 6.2  | 5.1  | 4.8  | 4.7  | 4.1  | 3.9  | 3.7  |
| Sand | Capillary potential  | 4    | 26   | 82   | 150  | 301  | 452  | 492  | 684  | 692  | 801  |
|      | Moisture content   | 39.6 | 34.2 | 30.2 | 22.8 | 19.4 | 17.5 | 16.4 | 15.5 | 15.0 | 14.6 |
| Loam | Capillary potential  | 5    | 29   | 55   | 144  | 269  | 371  | 489  | 556  | 715  | 796  |
|      | Moisture content   | 66.5 | 60.4 | 51.5 | 46.3 | 40.2 | 31.6 | 29.0 | 28.5 | 28.3 | 27.7 |
| Clay | Capillary potential  | 6    | 22   | 58   | 85   | 196  | 438  | 650  | 664  | 784  | 776  |

\* From "The Usefulness of Capillary Potential to Soil Moisture and Plant Investigators," L. A. Richards, *J. Agr. Research*, Vol. 37, No. 12, 1928, p. 732.

of the three soils with exertion of similar capillary potentials may be noted in the second vertical column. Likewise, it will be noted that, when capillary potentials of similar magnitude, namely, 492, 489, and 438, are exhibited by the sand, loam, and clay, respectively, the moisture percentages are 4.7, 16.4, and 31.6.

**Moisture Content of Soil.** It is a common practice to express the quantity of water in a soil in terms of percentage based on the weight of the soil. Frequently the weight of the moist soil is used as a base for calculation, as is the common practice in expressing the percentage of moisture in articles in trade, as hay, corn, vegetables, ores. The disadvantage of this method is that the base of calculation is different for each moisture content. To overcome this objection the weight of the dry soil (dried at 105°–110°C.) is often used as a base for expressing percentage moisture content in investigational work. Under this system the base is always the same regardless of moisture content. The following example will illustrate the difference between the two methods

of calculation. If to 100 grams of oven-dry soil 50 grams of water are added, 150 grams of moist soil are produced. With moist soil as a base, the soil contains  $50 \div 150 \times 100 = 33.3$  per cent moisture. With dry soil as a base, the soil contains  $50 \div 100 \times 100 = 50.0$  per cent moisture. It will be noted that at higher moisture contents there is a decided difference in the percentage moisture content as calculated by the two methods.

There is a valid objection to expressing the moisture content of soil as a weight percentage for practical purposes. In farming, soil is dealt with on a basis of depth, for example: A farmer plows to a certain depth or lays tile lines at given distances below the surface; plant roots penetrate the soil to different depths, depending on several conditions. Depth involves the concept of volume rather than of weight, as different soils have different volume weights. This is especially true of soils with high organic matter contents compared to those with little organic matter and also of well-granulated clays compared to sands. Frequently it is desirable to express moisture content on a volume basis as cubic inches of water per cubic foot of soil. In irrigation practice applied water is measured in inches of depth over an acre of surface, designated as *acre-inches*. Rainfall is also expressed in terms of inches in depth. If the moisture content of a soil is expressed in percentage based on the weight of the dry soil, the percentage of water by volume may be obtained by multiplying by volume weight. Table 17 shows the moisture content of an imaginary soil expressed by different methods.

TABLE 17

THE MOISTURE CONTENT OF A SOIL EXPRESSED BY DIFFERENT METHODS  
(The soil is hypothetical and assumed to have 43.4 per cent pore space with a volume weight of 1.50 and specific gravity of 2.65.)

| Percentage of Moisture Based on |          |                | Cubic<br>Inches of<br>Water per<br>Cubic Foot<br>of Soil | Pounds of Water per |             | Inches of<br>Water for<br>3-Foot<br>Depth |
|---------------------------------|----------|----------------|--|---------------------|-------------|---|
| Moist soil                      | Dry soil | Soil<br>volume |  | Cubic foot          | Acre-foot   |   |
| 20.0                            | 25.0     | 37.5           | 648.00   | 23.43               | 1,020,697.9 | 13.49                                     |

**Determination of Moisture Content.** The common method of determining the moisture content of soil is to weigh a sample of the moist

soil, dry it at a temperature of 105°–110°C., and reweigh when cool. Although this method is simple, time is required for the drying of the soil, samples of field soil must be taken to a laboratory where balances and an oven are available, and in studying moisture changes in field soils over a period of time repeated samplings are necessary, a condition which introduces an error due to lack of uniformity in the soil area. For years attempts have been made to devise a method for determining

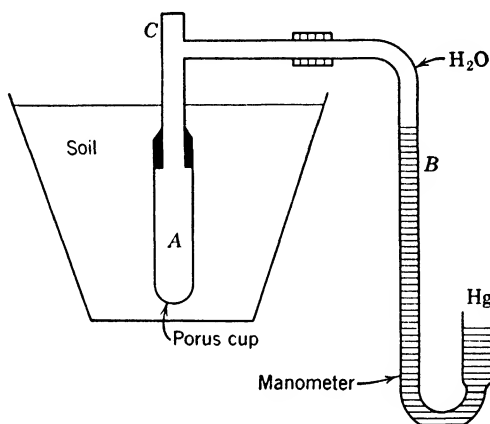


FIG. 41. A simple tensiometer for determining the tension-moisture curve. [From *Soil Physics*, by Baver, John Wiley & Sons, Inc., 2nd edition, 1948, p. 211—after Richards, *J. Agr. Research*, Vol. 37, pp. 719–742.]

soil moisture content quickly and without removing a sample. The conductance of soil between two electrodes placed in the ground and measured by some form of Wheatstone bridge has been studied. The conductivity would vary with moisture content, between certain limits, if no interfering factors were present. Changes in content of soluble salts, however, affect the conductivity to a greater extent than changes in moisture content, and so the method has not yet proved satisfactory. Bouyoucos and Mick have suggested the use of a plaster of paris block cast around two electrodes. The block can be buried at any desired depth, and insulated wires from the electrodes brought to the surface. The moisture content of the block quickly comes to equilibrium with that of the soil in which it is embedded, and, as the conductivity of the block is a function of its moisture content, readings may be quickly made by attaching the lead wires to a specially designed bridge.

More recently a unit composed of nylon fabric encased in perforated metal and containing two electrodes has been developed. This unit has

the advantage of registering moisture contents of the soil virtually to the saturation point while the gypsum block is not sensitive at moisture contents above field capacity.

Fletcher and Baver also have proposed new methods for measuring soil moisture content. Fletcher makes use of a specially constructed condenser which is placed in the soil, and, after equilibrium is attained with the soil moisture, readings are taken with a suitable capacitor. Baver suggests calculation of the moisture content from the heat conductivity of the soil.

Another device proposed by Richards and associates for measuring the movement of water in soils with moisture contents above the moisture equivalent is the tensiometer. This method is the same as that used for measuring capillary potential as described on pages 178 and 179.

**Classes of Retained Water.** The attraction between soil and water is so great that all the moisture is not removed by drying at a temperature of 110°C., although the quantity of moisture remaining is very small and a soil so dried is considered to be moisture-free for general purposes. If a thin layer of oven-dry soil is exposed in a desiccator over 3.3 per cent sulfuric acid, which will create an atmosphere with 98.2 per cent relative humidity, the soil will adsorb appreciable quantities of moisture. The percentage of moisture so adsorbed is designated the *hygroscopic coefficient* and will vary approximately with the colloidal content of the soil, although the nature of the colloidal minerals present has considerable influence on this value. Any amount of moisture a soil contains up to and including the hygroscopic coefficient is known as *hygroscopic moisture*. Water held in this condition does not move in the soil, is not used by plants, and in fact may exist in other than the liquid state.

Water held by a soil in excess of the hygroscopic coefficient is considered to exist in the form of thin films around soil particles and granules, in the larger pores of the granules, and in the angles or crevices made by the contact of particles. In general, this water may be thought of as a series of small wedges occupying the angles formed by particle contact and connected by films. This water is defined as capillary, from the old conception that the pore space in soil is similar to a series of capillary tubes containing moisture. Capillary moisture is susceptible to movement in the soil to a certain extent, may be absorbed by plant roots, and exists in the liquid state. The percentage of capillary water held by different soils varies with texture, structural conditions, organic

matter content, and the like. There has been much conjecture about the effect of organic matter on the capacity of soils to hold water and particularly water which is available for crop use. Information on this question was obtained by collecting samples of soil from fence rows and from adjoining fields which had been under cultivation for a long period of years. This procedure gave pairs of samples which were of similar texture but of different organic matter content. An average of the data for 25 pairs of samples is presented in Table 18.

TABLE 18

THE EFFECT OF ORGANIC MATTER CONTENT ON THE MOISTURE-RETAINING CAPACITY OF SOILS

| Sample                           | Organic Matter, per cent | Hygroscopic Moisture, per cent | Wilting Point, per cent | Moisture Equivalent, per cent | Available Moisture Capacity, per cent |
|----------------------------------|--------------------------|--------------------------------|-------------------------|-------------------------------|---------------------------------------|
| Fence row                        | 3.45                     | 1.51                           | 9.71                    | 24.93                         | 15.22                                 |
| Cropped                          | 2.09                     | 1.29                           | 9.08                    | 19.94                         | 10.85                                 |
| Difference due to organic matter | 1.36                     | 0.22                           | 0.63                    | 4.99                          | 4.37                                  |

An empirical method of measuring the power of different soils to hold moisture has been devised by submitting a thin layer of the soil to a pull of 1,000 times gravity in a porous cup placed in a specially constructed centrifuge. The percentage of water retained by a soil when submitted to this treatment is known as the *moisture equivalent*.

The percentage of moisture in a soil after drainage has ceased and, after capillary adjustments have taken place, is variously known as *normal field capacity*, *normal moisture capacity*, and *field-carrying capacity*. The value is also close to that of the moisture equivalent and is approximately two to three times the hygroscopic coefficient.

Hilgard proposed that the capacity of different soils to hold moisture might be compared by placing a short column of soil (about  $\frac{2}{5}$  inch) in a porous-bottomed cup. The cup is then set in a pan of water of such depth that the water surface touches the bottom of the soil column. When the soil becomes saturated, the cup of soil is removed from the pan and allowed to stand until drainage ceases. The percentage of water retained is called the *maximum moisture capacity*.

Table 19 shows the moisture equivalent, permanent wilting point, and the per cent of moisture retained by several western soils of different textures, as reported by Richards.

TABLE 19

THE INFLUENCE OF TEXTURE ON THE MOISTURE EQUIVALENT AND WILTING POINT AND THE CAPACITY OF SOIL TO RETAIN MOISTURE AGAINST GIVEN TENSIONS \*

| Type of Soil            | Moisture Equivalent, per cent | First Permanent Wilting, per cent | Moisture Retained under a Tension Equivalent to the Indicated Centimeters of Water, per cent |      |      |
|-------------------------|-------------------------------|-----------------------------------|--|------|------|
|                         |                               |                                   | 250  | 345  | 518  |
| Tujunga sand            | 2.6                           | 1.8                               | 2.3  | 2.1  | 2.1  |
| Holland sandy loam      | 6.9                           | 4.2                               | 7.6  | 6.4  | 5.5  |
| Romona fine sandy loam  | 9.2                           | 5.2                               | 10.1   | 8.3  | 7.8  |
| Greenfield loam         | 12.7                          | 6.3                               | 17.3   | 14.5 | 13.1 |
| Altamont clay loam      | 18.4                          | 10.0                              | 19.0   | 16.1 | 15.3 |
| Hanford silty clay loam | 24.4                          | 14.3                              | 29.5   | 24.3 | 22.8 |
| Yolo clay               | 45.9                          | 29.6                              | 52.3   | 45.1 | 43.5 |

\* From "Moisture Retention by Some Irrigated Soils as Related to Soil-Moisture Tension," L. A. Richards and L. R. Weaver, *J. Agr. Research*, Vol. 69, No. 6, 1944, p. 218.

**Movement of Soil Moisture.** Moisture may move in the soil in the liquid and in the vapor states. The movement of moisture in the soil in the liquid state takes place because of the unequal tensions developed under different degrees of curvature of the surface film; it is known as capillary movement. The cause of this movement may be explained as follows.

If a drop of water is placed on a glass plate covered with a film of grease, it will assume an almost spherical shape because the cohesion of the molecules tends to force the water into as small a volume as possible. The attraction between the grease and water offers small resistance to this tendency. A drop of water placed on soil particles has the same tendency to form a sphere, but then the cohesive force is largely overbalanced by the adhesion of the soil and water, and so the water is spread as a thin film over the soil particles. Now, if water occurs as wedges in the angles of adjoining soil particles with connecting films, as shown in Fig. 42, there is a tendency for it to form spherical droplets at each surface, as *A* and *B*. Droplet formation is prevented, however, because of the attraction of the soil and because the two surfaces are acting

against each other. Now, if the curvature of one film is greatly increased through the removal of moisture by a root, as is shown at *A*, its pulling power will be proportionately increased since the pull exerted by such a curved film is inversely proportional to the radius of curvature. As a result, water will be drawn toward *A* until the curvature of films *A* and *B* is equal. The equation for capillary movement may be written:

$p$  (pull) =  $2T/r$ , in which  $T$  is surface tension and  $r$  the radius of curvature. If it is assumed, for example, that the  $r$  of *A* is 1 and of *B* is 2, the pull exerted at the two points will be  $p = 2T/1$  compared to  $p = 2T/2$ . As the liquid in each case is water,  $T$  will be equal, and so the pull exerted by the film *A* will be twice as great as that exerted by film *B*.<sup>1</sup>

The distance and the rate of movement of capillary water have been the subjects of much discussion and investigation. It can be seen that, if the spaces between the soil particles are too large, the cohesion of the water molecules will be insufficient to raise it to any appreciable height; however, the movement will be rapid as friction will be small. This is the condition which exists in sandy soils. A somewhat analogous situation exists in the installation of telephone wires. If the posts are set too far apart, the wire will break because cohesion in the wire is insufficient to support the weight. On the other hand, in soils containing considerable silt and clay the spaces between particles will be small, and hence cohesion will support a relatively high column. Furthermore, in a fine-textured soil there is much more surface offering a greater opportunity for the action of the attractive force between soil and water which is the primary force holding the

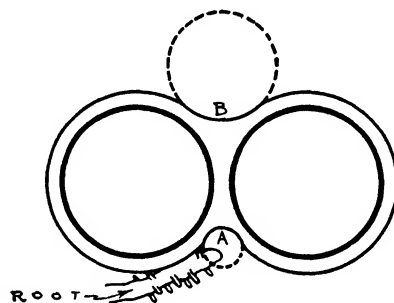


FIG. 42. As the root absorbs moisture from the accumulation between two soil particles, the film curvature increases, as is shown by the projected circles. As the force tending to draw water into a given portion of the film varies inversely with the radius of the curvature ( $P = 2T/r$ ), it follows that moisture will move to the feeding point of the root.

<sup>1</sup> Students particularly interested in capillary movement of soil moisture will find very intriguing the comparisons between this phenomenon and the flow of heat through a metal bar or the passage of an electric current through a wire.

water in the soil. Water may then be expected to rise more slowly but to greater heights in soils of a fine texture.

The principle may be illustrated by Fig. 43, in which a small and a large capillary tube are set in a basin of water.

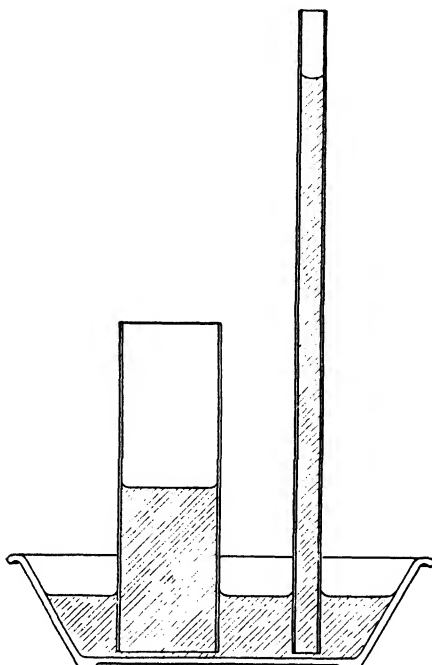


FIG. 43. Water rises to different heights in capillary tubes, depending on the diameters of the tubes. In small tubes there is a maximum of contact between the water and glass, giving a strong adhesive force, and the diameter is so small that cohesive forces can support a high water column. In a large tube adhesion between water and glass can be exerted on only a small part of the water, and the diameter is so large that cohesion can support only a short column.

The rate of rise of capillary water in a sand, silt loam, and clay loam is shown by the data in Table 20 together with the heights to which the water has risen after different time periods.

On a theoretical basis it has been calculated that moisture may be lifted from a free-water surface to a height of  $1\frac{1}{2}$  feet by coarse sand and to 150 feet by fine silt (particles from 0.01 to 0.002 mm. in diameter). However, these values have no significance from a practical viewpoint because the movement in the silt would be extremely slow, espe-

TABLE 20

## RATE AND HEIGHT OF CAPILLARY RISE OF WATER IN SOILS OF DIFFERENT TEXTURES \*

| Time     | Height of Rise, inches |           |      |
|----------|------------------------|-----------|------|
|          | Sand                   | Silt loam | Clay |
| ½ hour   | 13.5                   | 7.3       | 5.4  |
| 1 hour   | 14.3                   | 11.2      | 8.0  |
| 6 hours  | 16.6                   | 26.6      | 15.5 |
| 12 hours | 17.2                   | 35.3      | 18.5 |
| 1 day    | 18.4                   | 46.4      | 21.0 |
| 3 days   | 20.3                   | 65.4      | 24.7 |
| 6 days   | 21.8                   | 78.5      | 27.3 |
| 9 days   | 23.0                   | 86.3      | 28.8 |
| 18 days  | 25.3                   | 99.2      | 33.2 |

\* From *Soils and Fertilizers*, F. E. Bear, John Wiley & Sons, Inc., 3rd edition, 1942, p. 82. Used by permission.

cially after the water had risen a short distance, and because under field conditions capillary movement of moisture into the root zone of most crops does not take place from a free-water surface except when the water table is at no great depth.

The role of capillarity in supplying moisture to crops has undoubtedly been misinterpreted. Under field conditions moisture moves from soil of somewhat greater moisture content to soil containing less moisture. Seldom is the difference in moisture content very great, and as a result movement is slow—so slow that a plant will wilt before an appreciable quantity of water can move within reach of its roots from any great distance. Furthermore, as the supply of capillary water becomes less, movement is correspondingly slower. Unquestionably, however, considerable amounts of moisture move through short distances to the roots of growing crops, and hence capillarity is of some significance in supplying moisture to plants.

**Movement of Water Vapor.** The relative humidity of the air in soils containing appreciable quantities of moisture is approximately 100 per cent. Also, the pressure of the water vapor in the soil atmosphere increases with temperature. If, therefore, there is a considerable difference in the temperature of the soil at different depths, water

vapor may move from the warmer zone and condense in the cooler soil. It is conceivable that, if a long period of hot weather which has warmed the soil to a considerable depth is followed by a cold spell which cools the surface soil rapidly, appreciable quantities of water vapor may move up from the warm soil below and condense in the upper soil layer. Such movement may also occur in the fall when the lower soil horizons have not yet cooled to the temperature of the surface soil. Likewise, some moisture may condense in the soil surface from a warm atmosphere of high humidity.

### WATER IN RELATION TO PLANT GROWTH

Although the farmer considers soil moisture in relation to the fitting of a seedbed and to the tillage of his crops, and precipitation in relation to the loss of fertile surface soil through erosion, his primary interest in moisture is as a factor in crop growth. Not only is moisture necessary for the absorption of nutrients by plant roots, but also it is a nutrient itself and as such is as important as phosphate, potash, or nitrate. In fact, a growing plant contains a much larger proportion of water than of any other chemical compound. Plants are colloidal systems in which water serves as the liquid phase. The following questions emphasize pertinent points concerning the relation of water to plant growth.

#### Questions

1. How do plant roots absorb moisture?
2. How does water function in the absorption of nutrients by plants?
3. Is water effective in bringing nutrients within the reach of plant roots?
4. Is transpiration of service to plants?
5. How much of the retained water can plants absorb?
6. What is the wilting coefficient?

**Absorption of Moisture by Plants.** Moisture enters plant roots by the process of osmosis, which may be roughly defined as the movement of a liquid through a semi-permeable membrane caused by unequal concentrations on the two sides. It is customary to think of the cell sap within the root as having a greater concentration of soluble material than the soil moisture, and hence water passes in to equalize the concentration. A more correct view is to consider the concentration of water molecules in the cell sap reduced because of the quantity of soluble substances present, and hence the number of water molecules in the soil solution is greater. As a result, more water molecules strike

against a unit area of the exterior of the root membrane than against the interior, for the molecules in both cell sap and soil solution are in constant motion. As a result of the bombardment, water passes into the root from a zone of higher concentration of water to a zone of lower concentration.

If through any circumstance the concentration of soluble substances in the soil moisture exceeds that of the cell sap, the situation will be reversed and water will pass out of the root.

It should be recalled to mind that plant roots do not absorb the soil solution as such, as animals drink water containing soluble material. The water enters the roots as pure water without regard to the intake of any of the materials dissolved in it. The entrance of dissolved substances is entirely a separate process.

**The Role of Water in Nutrient Absorption.** It is generally believed that material must be in solution to pass through the plant-root membrane. Nutrients in the soil solution should therefore be in suitable condition for absorption. It must be pointed out, however, that the presence of a nutrient in the soil solution does not assure its use by the plant because the passage of ions and molecules through the absorbing membranes of the root is affected by various factors which cannot be discussed at this point. Furthermore, it cannot be assumed that nutrients must be in the soil solution to be used by plants. In the zone of intimate contact which exists between a root hair and a soil particle or a colloidal aggregate there is the possibility of the solution and absorption of nutrients. What proportion of the materials absorbed by plant roots comes from the soil solution and what directly from the particle surfaces it is impossible to tell.

**Movement of Nutrients in Soil Moisture.** Nutrients dissolved in the soil solution move with it, and so when moisture moves by capillarity to replace that which has been taken up by plants, a supply of nutrients may be moved near the roots. Although this action takes place through short distances only, the net result in the course of a growing season may add materially to the food supply of the crop. Vertical movement of soluble salts in capillary water has been observed more extensively than horizontal movement. During periods of drought a considerable accumulation of salts may develop at the soil surface. In some soils this may be sufficient to be visible, especially in regions of neutral or alkaline soils. The appearance of small quantities of salt on the surface of clods is a familiar sight even in humid regions. The movement of salts to the surface is much more pronounced in soils not occupied by growing plants because the roots absorb the moisture and

so reduce the quantity evaporated from the soil surface. Water from heavy rains has a tendency to displace the soil solution from the pore spaces and force it out of the soil into the drainage water. This process results in considerable loss of nutrients. There is also considerable mixing of the rain water with the soil solution, and a portion of the mixture drains away while the remainder constitutes the new soil solution which dissolves fresh supplies of nutrients from the soil complexes.

**Transpiration.** There has been much discussion concerning the value of transpiration to plants. Some cooling of the plant leaves may result from the evaporation of moisture from them, but there is a difference of opinion as to whether the plant is benefited by this effect. Some investigators think that transpiration is of assistance to plants in the movement of nutrients, particularly from the lower into the upper parts. Nutrients must be in solution for translocation to different parts of the plant, and, since water serves as the dissolving medium, it is of service in the growth processes aside from its role as a food material.

**Available Moisture.** As the quantity of moisture in the soil becomes less, the force by which it is retained increases. Finally, a point is reached at which the adhesion between soil and water equals the absorptive power of the plant roots, and hence no more moisture can be utilized by the plant. Theoretically this point represents the lower limit of available moisture. It varies greatly in soils according to the amount of surface exposed and therefore with the content of colloidal material. Since in the actual growth process it is impossible for plants to withdraw the last traces of available moisture from a soil, as shown in the discussion of *wilting point*, the quantity of water a given soil can retain in an unavailable state can only be determined by some laboratory method.

The upper limit of available water is assumed to correspond roughly with the percentage of water retained. On the other hand, as the last traces of gravitational water leave the soil slowly and the division point between gravitational and retained water may vary somewhat with the temperature, there is reason to assume that plants use some gravitational moisture. The quantities of available moisture which three soils of different texture will hold are shown in Table 21.

**Wilting Coefficient.** It is pointed out (p. 192) that the rate of transpiration of plants varies with environmental conditions which affect evaporation. The quantity of water lost by a given plant in a given time, say 1 hour, will also vary with the extent of the transpiring surface. When the quantity of water absorbed by the roots is less than the

TABLE 21

## AVAILABLE WATER HELD BY SOILS OF DIFFERENT TEXTURES \*

Water table at 6-foot depth. Moisture contents are for 3-foot depth of soil.  
Percentage moisture calculated on dry soil.

|                            | Dunkirk<br>Fine Sand | Wooster<br>Silt Loam | Brookston<br>Clay |
|----------------------------|----------------------|----------------------|-------------------|
| Water-retaining capacities | 6.5                  | 30.2                 | 30.5              |
| Wilting coefficients       | 3.5                  | 7.7                  | 17.5              |
| Available moisture         | 3.0                  | 22.5                 | 13.0              |

\* Data from *Soil Management*, F. E. Bear, John Wiley & Sons, Inc., 1st edition, 1924, p. 87. Used by permission.

quantity lost by transpiration, a negative moisture balance is set up in the plant, and very shortly the cells cease to be turgid and wilting sets in. Wilting will occur before a plant has taken all the moisture from a soil that it can, since the soil retains the last traces of available moisture with much force and the roots cannot take up moisture in the face of this resistance at a sufficiently rapid rate to equal transpiration. Furthermore, at this low moisture content capillary movement of moisture to the absorbing zone of the root is too slow to meet the demands of the plant. When a plant has wilted to such an extent that it will not revive when placed in a saturated atmosphere, it is said to be *permanently wilted*, and the percentage of moisture in the soil when permanent wilting occurs is designated the *wilting coefficient* or *wilting point* of that soil.

As is to be expected, the wilting point of soils varies greatly and is markedly influenced by the content of colloidal material and organic matter. Any given soil has about the same wilting coefficient for all common plants. The influence of soil texture on the percentage of moisture in soils when plants wilt is shown by the data in Table 22.

TABLE 22

## INFLUENCE OF SOIL TEXTURE ON THE WILTING COEFFICIENT

| Crop    | Coarse<br>Sand | Fine<br>Sand | Sandy<br>Loam | Loam | Clay<br>Loam |
|---------|----------------|--------------|---------------|------|--------------|
| Corn    | 1.07           | 3.1          | 6.5           | 9.9  | 15.5         |
| Sorghum | 0.94           | 3.6          | 5.9           | 10.0 | 14.1         |
| Wheat   | 0.88           | 3.3          | 6.3           | 10.3 | 14.5         |
| Pea     | 1.02           | 3.3          | 6.9           | 12.4 | 16.6         |
| Tomato  | 1.11           | 3.3          | 6.9           | 11.7 | 15.3         |
| Rice    | 0.96           | 2.7          | 5.6           | 10.1 | 13.0         |

### LOSS OF MOISTURE FROM THE SOIL

On all but a very small fraction of the farm land of the world, crops must draw their moisture from the supply stored in the soil, supplemented by rains. As the farmer sees his crops suffering from lack of water during periods of drought, he gives considerable thought to methods of increasing the moisture-retaining power of his soil and especially to ways of decreasing the loss of stored water. The conservation of soil moisture is uppermost in his mind as he cultivates his crops. A study of processes by which moisture is lost from the soil and of possible means of decreasing these losses is of practical significance. Questions which arise in a consideration of moisture loss from the soil are presented.

#### Questions

1. How much moisture is transpired by plants?
2. May the amount of transpired water be reduced?
3. What conditions affect evaporation of water from the soil?
4. How do mulches decrease evaporation?
5. What effect does soil cultivation have on evaporation of soil moisture?
6. Can losses of water by percolation be decreased?

**Loss by Transpiration.** As transpiration is simply an evaporation of moisture from plant surfaces, it is influenced by the same factors that affect the evaporation of water from any moist surface; exposure to direct sunlight, air temperature, humidity, wind movement, and atmospheric pressure are among the most important. To these must be added the characteristics of the plant itself, because transpiration is considerably more rapid from some plants than from others. This is due mainly to the number, size, and location of the stomata, although the nature and thickness of the leaf covering are also concerned.

Under uniform conditions a plant will transpire about the same quantity of water for each pound of dry plant material that develops. The weight of water in pounds transpired per pound of dry plant tissue produced above ground is known as the *transpiration ratio*. This ratio varies for different kinds of plants and for the same kind of plant grown under different climatic conditions. In some cases there is also a difference in transpiration ratio for different varieties of the same plant species, as is shown in Table 23. The great variation in water requirement of the same variety of plant, due to different climatic conditions prevailing in different years, is illustrated by the data in Table 24.

TABLE 23

## VARIATION IN WATER REQUIREMENTS OF DIFFERENT VARIETIES OF CROPS \*

| Pounds of Water Required on Basis of Pounds Produced of |       |             |           |       |             |                 |  |             |
|---|-------|-------------|-----------|-------|-------------|-----------------|--|-------------|
| Wheat   | Grain | Total plant | Oats      | Grain | Total plant | Corn            |  | Total plant |
| Turkey  | 995   | 364         | Canadian  | 1,416 | 399         | Iowa Silvermine |  | 302         |
| Kharkov   | 1,064 | 365         | Swedish   |       |             | Hopi            |  | 285         |
| Kubanka   | 1,111 | 394         | Select    | 1,103 | 423         | Northwestern    |  |             |
| Bluestem  | 1,573 | 451         | Burt      | 1,224 | 449         | Dent            |  | 280         |
|   |       |             | Sixty-day | 1,172 | 491         | China White     |  | 315         |

\* From "Relative Water Requirements of Plants," L. J. Briggs and H. L. Shantz, *J. Agr. Research*, Vol. 3, No. 1, 1914, pp. 8, 11, 17.

TABLE 24

## EFFECT OF SEASON ON WATER REQUIREMENT OF CROPS \*

| Pounds of Water Transpired per Pound of Dry Plant Tissue Produced in |                   |       |      |      |                    |         |
|--|-------------------|-------|------|------|--------------------|---------|
| Crop   | Variety           | 1911  | 1912 | 1913 | Greatest variation | Average |
| Wheat  | Kubanka           | 468   | 394  | 496  | 102                | 452.7   |
| Oats   | Swedish Select    | 615   | 423  | 617  | 194                | 551.7   |
| Corn   | Northwestern Dent | 368   | 280  | 399  | 119                | 349.0   |
| Sorghum  | Red Amber         | 298   | 237  | 296  | 61                 | 277.0   |
| Alfalfa  | Grimm             | 1,068 | 657  | 834  | 411                | 853.0   |

\* From Briggs and Shantz, *ibid.*, p. 56.

TABLE 25

WATER REQUIREMENT OF DIFFERENT CROPS AT AKRON, COLORADO<sup>1</sup>

| Average for 2 or 3 Years and Two or More Varieties |                                     |                |                                     |
|--|-------------------------------------|----------------|-------------------------------------|
| Crop   | Pounds water per pound plant tissue | Crop           | Pounds water per pound plant tissue |
| Millet   | 279                                 | Sugar beet †   | 397                                 |
| Sorghum  | 311                                 | Potato †       | 554                                 |
| Corn   | 375                                 | Sweet clover † | 770                                 |
| Wheat  | 532                                 | Alfalfa ‡      | 904                                 |
| Barley   | 534                                 | Oats           | 597                                 |

\* From Briggs and Shantz, *ibid.*, pp. 58-60.

† One variety for 2 years.

‡ Two selections of Grimm for 2 years.

In Table 25 are shown the transpiration ratios of several crops grown at Akron, Colorado. These results are much higher than would be expected for the same crops grown at lower altitudes and in regions of higher humidity with more cloudy days. Nevertheless, the data show that different crops vary greatly in their water requirement under similar climatic conditions.

**Reducing the Transpiration Ratio.** Alteration of environmental factors which affect evaporation will alter the transpiration ratio. Unfortunately, little can be done along this line except in the production of special crops on a limited scale. The erection of shades to protect plants from direct sunlight is practiced in the growing of special types of tobacco and medicinal herbs, such as ginseng. Dense windbreaks have some influence on transpiration for a limited distance. A high air humidity, obtained by sprinkling walks and floors of greenhouses, reduces transpiration. On a general scale, however, man can do little along this line.

It has been demonstrated that plants receiving an abundant supply of moisture transpire more freely than plants supplied with a limited quantity of water. Also, plants growing in a medium containing relatively small quantities of nutrients appear to transpire more moisture per pound of plant tissue produced than those growing in a medium containing an abundance of plant-food materials. Whether or not the

transpiration ratio of plants can be appreciably reduced under field conditions by applying liberal quantities of fertilizer has not yet been demonstrated.

**Evaporation of Soil Moisture.** Aside from the environmental factors which influence evaporation of moisture from any surface, previously mentioned, the following may affect loss of water from the soil surface. The shade afforded by growing crops and their protection of the soil from wind action are immeasurable but influential factors in reducing evaporation. Likewise, the higher humidity in the air in contact with the soil, resulting from the protection afforded by the crop, particularly by a low-growing, leafy crop, must have an influence on evaporation, especially in the morning and evening.

The nature of the soil surface is also significant. If the soil is loose so that air moves freely through it, evaporation will be increased. Unevenness or roughness of the soil that exists after cultivation or plowing also exposes more surface for evaporation, as do cracks in the soil. Cracking is especially prevalent in heavy soils during dry periods and results in much loss of water. A hard, compact surface in soils very high in clay may permit more capillary movement of moisture to the surface and therefore greater evaporation loss.

**The Function of Mulches.** A mulch may be roughly defined as any protective covering resting on the soil. It may be of entirely foreign material, such as shavings, gravel, boards, straw, paper, or it may be a layer of dry soil. Whatever it is, the chief function of the mulch is to protect the moist soil from the direct rays of the sun, and from direct contact with wind and warm dry air. Other effects of a mulch which sometimes result are the prevention of soil cracking through a reduction of evaporation, a limiting of capillary movement of moisture, and a greater absorption of rainfall through impeding surface runoff. Furthermore, a mulch prevents the "running together" of the surface soil under the impact of rain drops and the development of a structural condition which restricts the intake of water. It has also been shown that mulches of peat sometimes reduce the intake of moisture by absorbing the rain and later losing it through evaporation. A mulch of gravel, on the other hand, permits ready absorption of rain by the soil.

**Cultivation and Loss of Water by Evaporation.** The stirring of the moist surface soil with tillage implements results in increased aeration and exposure of more surface to wind and sun, all of which hastens drying. A layer of dry soil is thus produced which serves as a mulch over the moist soil beneath. This mulch functions like any other in reducing evaporation and in moisture conservation. Also, cultivation

during dry periods reduces loss of moisture by filling cracks and keeps the soil in a suitable physical condition for the absorption of rainfall. In soils very rich in clay, cultivation may interrupt the movement of capillary moisture to the surface and so prevent excessive loss of soil water.

In the great majority of soils, however, the rate of evaporation from the surface is more rapid than the capillary movement of moisture, and hence a layer of dry soil (constituting a mulch) is developed without the aid of cultivation. The loss of moisture during the development of this natural mulch appears to be no greater than that during the formation of a mulch through tillage, and there seems little difference in the effectiveness of the two in preventing further moisture losses. In this connection it should be mentioned that Veihmeyer has shown that, before the soil has become dry enough to cultivate into a fine mulch, most of the moisture which will be lost has already evaporated. Numerous experiments have been conducted to determine the effect of cultivation on the yield of crops. Most of them have been carried out on soils which do not tend to pack or cake but to retain a good state of tilth either because of a moderate clay content or of a relatively high organic matter content. In Table 26 are presented sample results from several of these experiments.

TABLE 26

COMPARATIVE EFFECT OF REMOVAL OF WEEDS WITHOUT DISTURBING THE SOIL,  
NORMAL CULTIVATION, AND WEED GROWTH ON THE YIELD OF CORN

| State    | Soil Type                 | Duration of Experiment, Years | Soil Treatment and Yield, bushels |                               |
|----------|---------------------------|-------------------------------|-----------------------------------|-------------------------------|
|          |                           |                               | Scraped,                          | Cultivated                    |
| Ohio     | Brookston silty clay loam | 6                             | 72.3                              | 3 times, 69.8                 |
| Ohio     | Miami silty clay loam     | 6                             | 56.3                              | 3 times, 59.7                 |
| Illinois | Brown silt loam           | 6                             | None—weeds allowed to grow, 7.0   | Cultivated with blades, 53.0  |
| Illinois | Brown silt loam           | 6                             | 53.3                              | Cultivated with shovels, 51.1 |

It is well recognized that utilization of water by weeds is one of the greatest sources of soil-moisture loss. Tillage is the most practical method of weed control, and hence the tendency of soils to mulch themselves does not eliminate the need for careful cultivation of the soil in

the preparation of a good seedbed and for adequate tillage of row crops during the growing season. However, if the soil is comparatively free of weed seed and care is taken to stimulate the germination of as many weed seeds as possible during seedbed preparation, the number of cultivations of the growing crop may be reduced. Much progress has been made in the control of weeds through use of chemicals. Perhaps it is not



FIG. 44. Lister furrows on the contour not only prevent erosion but also hold much water until it can be absorbed into the soil, as is taking place on this Oklahoma field. [Courtesy of Soil Conservation Service.]

too optimistic a view to assume that eventually a smaller number of cultivations may be required to control weeds in certain crops.

In heavy soils which do not maintain a good state of tilth but easily become compact, cultivation may be of value from the standpoint of increased aeration. Baver cites an example from the results of Ohio experiments, as follows: Corn on Paulding clay yielded 27.7 bushels of corn in 1927 when weeds were removed by scraping without stirring the soil, and 41.4 bushels with three normal cultivations.

**Percolation Losses.** Traveling eastward from the arid areas, one passes through the Western Plains region where only occasionally is rainfall sufficient to wet the soil to the depth of the permanent water

table. As the climate becomes more humid, losses of water by percolation assume proportions of practical significance. Distribution of rainfall is probably as important as total precipitation in inducing percolation, for a moderate amount of rain falling in the course of a few weeks may result in more drainage water than a much larger amount distributed throughout the year. Some other factors affecting percolation are (1) soil texture, (2) soil structure, (3) depth of soil and nature of underlying material, (4) loss by evaporation and transpiration, and (5) organic matter content of the soil. In some of the sandy soils in areas with long warm seasons and high rainfall the quantity of moisture percolating through the soil is unbelievably large.

Unfortunately, little can be done to reduce the loss of water through percolation. Increase in moisture-retaining power of the soil by inducing granulation of heavy soils and by compaction of sandy soil, coupled with an increase in organic matter content, is about all that can be suggested.

#### RUNOFF WATER

An appreciable proportion of the precipitation does not enter the soil but runs off over the surface in virtually all climatic zones regardless of how small the total precipitation may be. The damage done through this process of runoff of water is so widespread that the problem is one of universal concern, and as a result much attention has been given to methods of controlling runoff, particularly within the last twenty years. This subject is covered in considerable detail in Chapter 16.

# 9

## Soil Organisms— Kinds, Numbers, Activities, and Relation to Soil Productivity

The soil is the home of innumerable forms of plant and animal life that range in size from those too small to be seen with a powerful microscope to larger forms such as earthworms. Through a great variety of activities these organisms contribute to the productive capacity of the soil; for example, they are intimately concerned with the cycles or series of transformations of plant-food elements from one form to another. A better appreciation of the importance of living organisms in soil development and productivity comes through a knowledge of the general nature of the soil population, of the specific functions of the various forms, and of the influence of environmental factors on their survival and activities. A stimulation of desirable soil microbial activities can be obtained through the correction of unfavorable soil conditions by such practices as liming, drainage, adding organic matter, proper tillage operations, using good cropping systems, controlling parasites, and inoculation. The study of organisms living in the soil and the part they play in plant growth may be undertaken under four headings.

### Objectives

- A. The general nature and extent of the soil population.
- B. Activities of soil microbes in relation to the growth of higher plants.
- C. The role of microorganisms in the development of soils.
- D. Relationships between higher plants and soil microorganisms and among microorganisms themselves.

### NATURE AND EXTENT OF THE SOIL POPULATION

The original concept of the soil population as predominantly bacterial has been expanded through years of investigation to include also the numerous groups of fungi, actinomycetes, protozoa, algae, and many small

invertebrate animals. Carefully conducted studies have shown that soil characteristics such as structure, aeration, drainage, organic matter content, and reaction, as well as tillage practices and kinds of crops grown, exert pronounced effects on the abundance of the soil organisms as a whole and on the relative numbers of the various species. A general knowledge of the nature of this population should be of considerable interest to all who are concerned with the use of the land because we owe the continued development of higher plants to the activities of these tiny living things. A few questions will serve as guides in the study of this subject.

### Questions

1. What are the kinds, numbers, and characteristics of soil organisms?
2. What are the nutritional requirements of soil organisms?
3. How do the environmental factors of temperature, moisture, acidity, aeration, salts, light, and organic matter affect the nature of the soil population, and can these factors, in any way, be controlled by man?
4. What factors affect the distribution of microbes through the soil?

**Kinds of Organisms Found in Soils.** If we were to consider all life in or springing from the soil, it would be necessary to include all plants, ranging from large trees to the smallest microbes, with an almost indefinite number of life forms between the two extremes in size. In our discussion, however, we shall direct our attention to the bacteria, fungi, actinomycetes, and algae belonging to the plant kingdom; and to the protozoans, nematodes, and a few of the larger animals that live in the soil. A summary of the more important groups of soil organisms is presented in Chart 6.

**Bacteria.** In cultivated soils bacteria usually exceed all other organisms in numbers and kinds. They are single-celled plants, the larger individuals seldom exceeding 0.005 mm. in diameter. Generally they are regarded as the simplest and smallest forms of life. Most of them fall within the colloidal-particle size range. The numbers of bacteria present in soils vary greatly since many conditions affect their growth and since they can multiply in numbers with extreme rapidity by elongating and dividing. From each individual a new one may be formed in less than 20 minutes, and hence sudden increases in numbers may occur when soil conditions are favorable. Likewise, if soil conditions become unfavorable a rapid decrease in numbers may result.

The numbers of bacteria in soil, as determined by the plate method, vary widely, generally between 0.3 million and 95 millions per gram of soil. On the other hand, by the aid of microscopic methods the numbers

**Chart 6. Summary of the More Important Groups of Soil Organisms****PLANTS****I. Bacteria****A. Heterotrophic**

- 1. Nitrogen fixers
  - (a) Symbiotic
  - (b) Non-symbiotic
    - (1) Aerobic
    - (2) Anaerobic
- 2. Those requiring fixed nitrogen
  - (a) Spore formers
    - (1) Aerobic
    - (2) Anaerobic
  - (b) Non-spore formers
    - (1) Aerobic
    - (2) Anaerobic

**B. Autotrophic**

- 1. Nitrite formers
- 2. Nitrate formers
- 3. Sulfur oxidizers
- 4. Iron oxidizers
- 5. Those that act on hydrogen and various hydrogen compounds

**II. Fungi**

- A. Yeasts and yeast-like fungi
- B. Molds
- C. Mushrooms

**III. Actinomyces****IV. Algae**

- A. Blue-green
- B. Grass-green
- C. Diatoms

**ANIMALS****I. Protozoa**

- A. Ciliates
- B. Flagellates
- C. Amoeba

**II. Nematoda**

- A. Those that feed on decaying organic matter
- B. Those that feed on earthworms, protozoa, bacteria, etc.
- C. Those that infest the roots of higher plants

**III. Earthworms \*****IV. Other larger animals (worms, insects, ants, snails, spiders, mites, rodents, millipedes, centipedes, etc.) \***

\* These are frequently referred to as macroorganisms; the others in this chart as microorganisms.

have been found to range from 1 billion to 4 billions per gram of soil. It is not unlikely that the total weight of bacterial substance per acre to a depth of 7 inches of soil exceeds 500 pounds (live weight) in good soils.

Soil bacteria may be divided broadly into two large groups, based on their energy requirements: (1) the heterotrophic bacteria, which obtain their energy and carbon from complex organic substances; (2) the autotrophic bacteria, which can obtain their energy from the oxidation of inorganic elements or compounds, their carbon from carbon dioxide, and their nitrogen and other minerals from inorganic compounds.

In the latter group are found such organisms as the nitrite formers, the nitrate formers, the sulfur-oxidizing bacteria, the iron oxidizers, and those that act on hydrogen and its compounds.

The heterotrophic group may be subdivided into two groups: (1) the nitrogen-fixing bacteria, and (2) those that require fixed nitrogen. There are two groups of nitrogen-fixing bacteria, the symbiotic and non-symbiotic, which are discussed in more detail elsewhere (pp. 216-222). The non-symbiotic group may be subdivided further into aerobes and anaerobes. The former require an abundant supply of oxygen, and the latter function in the absence of oxygen. There are two groups of bacteria requiring fixed nitrogen. They are the spore formers and non-spore formers, and each of these groups may be further divided into aerobic and anaerobic forms.

These numerous soil bacteria differ considerably in their nutrition and in their response to environmental conditions. Consequently, the nature and abundance of the various types of bacteria depend both on the available nutrients present and on the soil environmental conditions.

*Fungi.* On the basis of their morphology, fungi may be classified into three groups: (1) yeasts and yeast-like fungi, (2) molds, including the various filamentous fungi, and (3) the mushroom fungi. These organisms vary greatly in structure and size from the simplest yeasts and molds to the more complex mushroom forms. They are all devoid of chlorophyll and must therefore obtain their energy and carbon from complex organic substances.

The members of the first group are found in the soil only to a limited extent and are believed to be of no great importance in soil development or productivity.

The other two groups normally are found in great numbers, with the mushroom fungi being particularly abundant in forest soils. The mushroom fungi form a very extensive mycelium, sometimes producing fruit-

ing bodies in the form of mushrooms, puffballs, and toadstools. Some of the fungi of this group produce an associative growth with the roots of higher plants, especially forest trees, often referred to as fungus rot or mycorrhiza.

In this relationship it is believed that the fungi in some way aid the higher plants in obtaining their nutrients and that the fungi obtain food and energy from the roots of the trees. The soil fungi are usually outnumbered by bacteria except in acid soils, in soils that have been heavily manured, or in those that are naturally rich in organic matter. Fungi respond especially well to aeration and can make extensive growth only under aerobic conditions. They make a vigorous growth in acid, neutral, or alkaline soils, and many fungi are favored by the lower *pH* values.

The number of fungi ordinarily found per gram of soil has been from 8,000 to somewhat over 1,000,000. This would probably be the equivalent of 1,000 to 1,500 pounds per acre (7 inches deep). It is observed, therefore, that fungi may be outnumbered by bacteria but that the fungi present a greater mass of growth per unit volume of soil. This is due, of course, to the much greater size of a fungus mycelium in comparison to a bacterial cell.

*Actinomyces*. The actinomycetes may be looked upon as occupying a position, from the morphological point of view, between the bacteria and the fungi. They are frequently spoken of as *ray fungi* or *thread bacteria*. The actinomycetes resemble bacteria in that they are unicellular and of about the same size in cross section. They resemble the filamentous fungi in that they produce a very extensive and profusely branched filamentous network. Many of these organisms reproduce by means of spores, and these spores appear very much like bacterial cells.

These organisms are present in great abundance in soil, making up as much as 60 per cent of the colonies that develop on plates containing artificial media inoculated with a soil extract. The numbers of actinomycetes may vary between 0.1 million and 36 millions per gram of soil. In actual weight of live substance per acre, they may exceed bacteria but as a rule will not equal fungus tissue. Under optimum soil conditions the total weight of actinomycetes on the average may exceed 700 pounds per acre of soil (7 inches deep).

For the most part, actinomycetes are aerobic, and in comparison to most bacteria they are more sensitive to changes in reaction and are active over a narrower range of acidity and alkalinity. The optimum development of actinomycetes occurs between *pH* 6.0 and *pH* 8.0, and practically no growth takes place below *pH* 5.0 with most species.

They make more extensive development than bacteria in soils of low moisture content, although the actinomycetes develop well in fairly moist soil under aerobic conditions. These organisms are especially numerous in soils of high organic content which are not too strongly acid. The odor of freshly plowed land noticeable at times is believed to be due to the products of the activities of actinomycetes.

Actinomycetes perform a very important function in soils by breaking down organic matter and setting free the plant nutrients it contains. Apparently these organisms are able to attack humus which would be very slowly decomposed by bacteria, and so release nitrogen which might long remain unavailable to higher plants. The actinomycetes thus perform a most important function, from the standpoint of soil fertility, in helping keep the soil nitrogen in circulation.

*Algae.* The chlorophyll-bearing microscopic plants are called algae, and they form a very extensive and important group of soil organisms. They are universally distributed in the surface layers of soil wherever moisture and light are available. The humidity of the soil appears to be a very important ecological factor in their distribution, for moist soils contain many more species than dry soils, although they are able to retain their vitality for long periods even after prolonged drought.

Algae obtain nitrogen and minerals from the soil, carbon from the carbon dioxide of the atmosphere, and utilize the energy of the sun as long as they have free access to light. Below the soil surface, in the absence of light, they act in a manner similar to fungi in that they utilize the energy derived from the decomposition of organic materials. Algae may thus live and function much like higher plants, or they may perform like the lower forms of plant life.

The algae vary greatly in size and shape; but the species commonly found in soils are mostly microscopic in size and are either unicellular or filamentous in structure.

They are usually classified into the following three groups: (1) blue-green, (2) grass-green, and (3) diatoms. Members of the first group frequently contain pigments other than chlorophyll and may thus vary in color from blue-green to violet or brown. The second group usually contains only chlorophyll, but occasionally other pigments may give them a yellow-green color. Members of the last group are golden brown owing to a blending of chlorophyll with other pigments. In the absence of light the chlorophyll in any species may or may not be retained.

As would be expected, algae are most abundant in the upper soil horizons where environmental conditions are most favorable for their development. Members of the blue-green and grass-green groups usu-

ally outnumber the diatoms. It is estimated that a soil under favorable conditions may contain 100,000 of each of either the blue-green or grass-green algae per gram. Diatoms usually are quite numerous in old garden soils.

The development of algae in the soil results in an increase in the supply of organic matter and in transforming, temporarily, soluble forms of nitrogen and minerals into organic or insoluble forms. By so doing they aid in reducing the leaching losses of soluble nutrients, but during the cropping season this may result in competition with higher plants. Furthermore, algae aid in the decomposition of organic material and exert a solvent action on certain rocks and minerals and thus affect the formation of soils. They appear to encourage the fixation of nitrogen by living symbiotically with *Azotobacter*, and perhaps the blue-green algae are able to acquire some free nitrogen themselves. It has been suggested that algae, by taking in carbon dioxide and giving off oxygen, aid rice plants growing in swamp soils. It is impossible to state specifically or with any degree of certainty the role played by algae in the various soil processes.

*Protozoa.* Protozoa are generally accepted as the simplest form of life belonging to the animal group. They are all microscopic in size and unicellular but larger than bacteria and more complex in their activities. In general, cells of protozoa exhibit a much higher development than bacteria. These organisms may move in the soil by means of cilia,<sup>1</sup> flagella, or pseudopodia, and this difference in the nature of their locomotion offers a convenient basis for classifying them. Thus they may be grouped as follows: (1) ciliates, (2) flagellates, and (3) amoeba. All three groups are abundantly represented in the soil, members of the last two groups being especially abundant. It is reported that soils may contain 500,000 to 1,000,000 flagellates, 100,000 to 500,000 amoeba, and 80 to 1,000 ciliates per gram. They might amount to a total weight of 200 or 300 pounds to the acre.

Because it is difficult to free protozoa from bacteria, the exact nature of the physiology of these small animals is still a matter of conjecture. Their nutritional habits are not well known, although it is certain that in the soil they depend on the organic matter as a source of food. Physiologically, however, protozoa vary considerably. Certain groups may ingest bacteria and thus indirectly affect the growth of higher plants. Several years ago a theory regarding soil productivity was advanced

<sup>1</sup> Cilia are hair-like projections capable of vibratory motion; flagella are whip-like appendages; and pseudopodia are temporary protrusions of the cell and may be used for locomotion.

which was based on the interrelationships of protozoa and bacteria. The protozoa feeding on bacteria were believed to decrease soil fertility because the bacteria at that time were looked upon as the only organisms capable of decomposing organic matter. Obviously this theory does not hold because many types of organisms are now known to have the capacity for breaking down organic matter. Granting that protozoa may feed on bacteria, their effect will be either beneficial or harmful, depending on the kind of bacteria consumed. By feeding on the nitrifying bacteria, protozoa would prove detrimental, but decidedly beneficial if they aided in destroying disease-producing bacteria. Under certain special conditions, as in waterlogged soils or in the presence of an abundance of organic matter, protozoa may keep the bacterial flora as a whole in check. It is believed, however, that the results of the relationship between protozoa and bacteria are largely beneficial because protozoa feed on pathogenic bacteria and keep important groups of bacteria in a youthful state for a longer period of time (by consuming the older bacterial cells), thus stimulating specific bacterial processes.

*Nematoda.* Among the various worms found in the soil, nematodes occupy a prominent position. Billions of them are found in each acre of soil, and on the basis of their food requirements three groups are distinguished: (1) those that feed on decaying organic matter; (2) those that feed on earthworms, other nematodes, plant parasites, bacteria, protozoa, and the like; and (3) those that infest the roots of higher plants, passing a part of their life cycle embedded therein. Nematodes are sometimes called eel worms; they are round or spindle-shaped and usually have a pointed posterior. Most nematodes are microscopic in size.

Members of the first group are in greater abundance in most soils than those of the other two groups, although the last group is the most important from the agricultural point of view. The roots of a great number of plants, such as the English pea, cowpea, tomato, and carrot, are entered by certain species of the third group, and a great deal of damage is often done. Investigations are indicating that nematode damage is much more extensive than originally thought. Nematodes may also become serious pests in greenhouse soils unless special care is taken to avoid infestation. Not only do the nematodes injure the plant roots themselves, but also by puncturing the plant they prepare an entrance for other parasites. Not all the activities of nematodes are harmful to the growth of higher plants, for they aid in bringing about an intimate mixture of the mineral and organic matter in soil and in breaking down organic materials. They may also improve soil aeration.

*Earthworms.* Perhaps the most important group of the larger animals inhabiting the soil is the common earthworm, of which there are several species. These organisms prefer a moist environment with an abundance of organic matter and a plentiful supply of available calcium. Consequently, earthworms are found most abundantly, as a rule, in heavy soils which are high in organic matter and are not strongly acid and occur only sparingly in acid sandy soils low in organic matter.

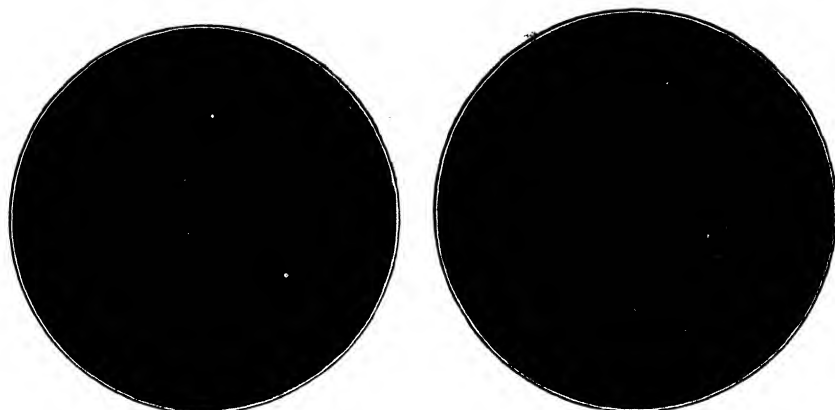


FIG. 45. Clay loam soil before (left) and after (right) being thoroughly worked by earthworms. Note the almost complete granulation resulting from passage of the soil through the digestive tract of the worms.

Obviously, then, the number and activity of the earthworms vary greatly from one location to another and, as with other soil organisms, figures indicating numbers are merely suggestive. The number of earthworms in the plowed layer of an acre may range from a few hundred or even less to more than a million. It has been estimated that between 200 and 1,000 pounds of earthworms are present in an acre of soil.

It is believed that in some soils these organisms may pass several tons of soil through their bodies annually and in so doing bring about an increased availability of plant nutrients. The large number of little mounds of earthworm casts are very noticeable in the early spring on soil sparsely covered with grass. Plant nutrients contained in both the mineral and organic portions of the soil are released, the nitrogen of the organic matter being affected particularly. Holes left in the soil by earthworms increase soil aeration and drainage. Furthermore, these organisms bring appreciable quantities of soil from the lower to the upper horizons, resulting in considerable soil mixing. Earthworms may drag undecomposed organic matter into the soil to serve as food and

protection. This is particularly noticeable in uncultivated soils, especially in certain forest soils. The development of the so-called mull layer (*A* horizon) in certain forest soils is due to this type of earthworm activity.

Earthworms are often objectionable when present in great abundance in lawns and especially in the soil of golf greens. Heavy applications of organic manures tend to increase the number present.

*Other Larger Animals.* In addition to earthworms, other groups of the larger animals inhabit the soil, namely rodents, ants, snails, spiders, mites, millipedes, centipedes, and various other worms and insects. Some of these organisms may spend all, and others only a part, of their life cycle in the soil.

The effects of these animals on the soil are beneficial for the most part. As a result of their activities (burrowing habits, for example), considerable quantities of soil are transferred and some disintegration of soil particles occurs. Their activities tend to increase aeration and improve the drainage of soils. Although soils may be directly benefited by their activities, it is obvious that they may prove unfavorable to agriculture under certain conditions.

Some of these animals cause various chemical changes in the soil either directly by their digestive processes or indirectly by influencing the activities of the soil bacteria, fungi, and other microscopic organisms. They may feed also on certain other soil organisms like algae, fungi, and protozoa. Much damage is done to certain crops by representatives of these groups, especially by some of the insects. From these few remarks it is evident that the animal population contributes to the complex system of activities going on in the soil.

**Nutrient Requirements of Soil Organisms.** Soil organisms have, in general, the same nutrient-element requirements as higher forms of life. For their growth and development they all require supplies of energy in addition to the several essential elements, including carbon, hydrogen, oxygen, nitrogen, phosphorus, potassium, sulfur. With the exception of algae all the important soil microbes are devoid of chlorophyll, and they must obtain their energy either from the oxidation of simple inorganic substances, as the autotrophic bacteria do, or from complex organic substances, as do most bacteria (heterotrophic organisms), all the fungi, and protozoa. Most soil organisms are thus dependent on the organic matter of the soil for their supply of food and energy. The autotrophic bacteria and algae obtain their carbon from carbon dioxide. Oxygen and compounds containing hydrogen are usually present in soils in sufficient quantities, although the former may be

deficient in poorly drained soils, and under such conditions certain microbes get their oxygen from inorganic compounds such as nitrates or sulfates. Nitrogen is one of the most important elements in the nutrition of microbes and may be used in the form of complex organic substances or in simple inorganic compounds, like ammonium or nitrate salts. Certain groups of bacteria have the ability to utilize the free nitrogen of the air and build it into their protoplasm, thereby increasing the soil's supply of combined nitrogen. Microbes obtain the other necessary mineral elements from the soluble salts in the soil and from the ash of decaying organic matter.

**The Influence of Soil Conditions on Microorganisms.** As has been indicated, the environmental conditions determine the nature of the microbial population present at any given time in the soil. In general, the fertile, heavy soils rich in organic matter contain many more microbes than the light soils poor in organic matter. If we examine the changes which occur in the microbial population of soils, we find certain causes for these variations. Let us consider briefly some of these causes.

*Temperature.* The optimum temperatures for the growth of most soil microbes are considerably higher, as a rule, than those which prevail in the soil even in summer. Consequently, it may be assumed that microorganisms never reach their highest level of activity in soil, and thus they utilize only a part of the potential energy sources. Temperature regulates the reacting velocities of chemical and biological changes occurring in the soil. Within a rather narrow range the rate of biological reactions increases two to three times for each increase in temperature of 10°C. Roughly, the limits of microbiological functions are reached with a temperature of 80°C. For the majority of the soil organisms the optimum temperature is about 35°C., although they can grow at rather wide temperature ranges and may adapt themselves readily to gradual changes in temperature.

*Moisture.* Another major factor affecting the numbers and activities of soil microorganisms is soil moisture. The influence of moisture depends to a large extent on the nature of the soil and the nature of the organisms concerned. The optimum amount of water for most soil organisms is between 50 and 70 per cent of the water-holding capacity of the soil, about the same as for most higher plants. Most soil organisms have the ability to withstand rather wide extremes in soil-moisture content, thus insuring their wide distribution in soils in spite of periodic changes in the moisture supply.

*Acidity.* The degree of acidity or alkalinity of the soil is of particular importance in influencing the activities and relative abundance of the

different groups of soil organisms. It is frequently noted that the proportion of fungi to bacteria and actinomycetes is greater in acid than in neutral soils; thus it appears that an acid soil favors the development of fungi but is unfavorable to the development of other forms. Usually the beneficial organisms function best in a soil that is approximately neutral in reaction. As a rule, the actinomycetes prefer a reaction of 7.0 to 7.5, the bacteria and protozoa from 6.0 to 8.0, and the fungi from 4.0 to 5.0. Below pH 6.0 azotobacter, in general, are inactive. In strongly acid soils legume bacteria fail to develop and function normally, and as a result poor inoculation is obtained and the organisms do not persist in the soil for any great length of time. The nitrifying organisms are also sensitive to a highly acid condition. Thus it becomes evident that the optimum in acidity for the majority of the soil population (especially the more desirable groups) is essentially the same as for most higher plants. The tolerance of soil organisms for acidity, as of higher plants, is influenced considerably by other conditions like nutrient supply and favorable moisture content and temperature.

*Aeration.* The development and activities of soil organisms are greatly affected by the concentration and rate of supply of certain gases (particularly oxygen, carbon dioxide, and nitrogen) in the air. Oxygen is needed for oxidation processes, carbon dioxide as a source of carbon for autotrophic organisms, and nitrogen for the nitrogen-fixing organisms. Abundant oxygen favors the activities of the nitrite and nitrate formers, the nitrogen fixers, fungi, actinomycetes, and other organisms which oxidize organic matter. Light soils frequently are excessively aerated, and this condition restricts the accumulation of organic matter, whereas heavy soils frequently are insufficiently aerated. Poor aeration favors reduction processes. Soil aeration is governed primarily by fluctuations in soil moisture. Aeration increases with a decrease in soil moisture, whereas an excess of water tends to encourage anaerobic conditions.

*Salts.* The addition of mineral salts to the soil influences the nature and activities of the soil population in several ways. They may stimulate the growth of higher plants, giving greater crop residues, thereby increasing the available energy supply, and resulting in increased microbial activities. With this increase in available energy, the supply of mineral elements may then become limiting factors. This is especially true of nitrogen, phosphorus, potassium, calcium, and magnesium. The addition of mineral nutrients tends to produce a more desirable balance in the concentration of the soil solution for microbial activities, although an excess of soluble salts may prove injurious. The action of the

various mineral salts depends on the nature of the salt, type of soil, and the nature of the organism concerned.

**Light.** Direct sunlight is highly injurious to most forms of soil micro-organisms, and many are instantly killed when exposed to it. Diffused daylight appears to have an inhibiting effect on most bacteria although it has little effect on the growth of fungi. The growth of algae, however, is stimulated by diffused daylight, but direct sunlight appears to be detrimental.

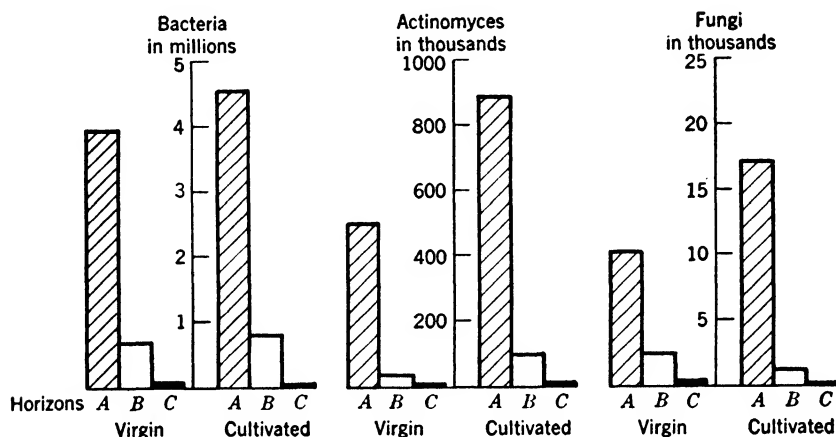


FIG. 46. Vertical distribution of microorganisms in a cultivated and virgin Carrington silt loam. [Data from *Iowa Research Bull.* 132.] The values for the A horizons are the averages for the  $A_1$ ,  $A_2$ , and  $A_3$  horizons. The values for the C horizon of the cultivated soil are the averages for the  $C_1$  and  $C_2$  horizons. All values refer to the number of organisms per gram of air-dry soil.

**Organic Matter.** Among the various factors affecting soil organisms, the influence of organic matter is certainly one of the most important, especially under humid conditions. Since organic matter is the source of food and energy for the majority of the organisms, obviously those soils abundantly supplied with this material are capable of supporting a more dense microbial population than those low in organic matter. There is also an influence of organic matter on the structure, aeration, water-holding capacity, and temperature relations of soils.

**Distribution of Microbes in the Soil Profile.** The greatest number of microbes as a rule occurs in the surface layers (A horizon) of the soil, as shown in Fig. 46, although the number at the very surface of cultivated soils may be relatively low owing to a lack of moisture and

to the germicidal action of the sunlight. In forest and meadow soils the greatest number of individuals and varieties of microbes is at or very near the surface of the soil. The number in all soils decreases with depth, below the *A* horizon, and the rapidity of this decrease in numbers varies with the soil conditions, especially the distribution of organic matter, aeration, and acidity.

The rate of decrease is relatively slow in most arid and semi-arid soils. The microbial population, therefore, is densest in that part of the soil profile where the food supply is plentiful and where the environmental conditions are most favorable, and this condition usually is found at or near the top of the soil profile.

### ACTIVITIES OF SOIL MICROBES IN RELATION TO THE GROWTH OF HIGHER PLANTS

The chemical processes that take place within the soil are, for the most part, dependent on the activities of living organisms. The growth and activities of microbes are intimately concerned with the availability of nutrients, and hence the existence of higher plants depends on the activities of soil microbes. After all, a productive soil is characterized not necessarily by the mere presence of large quantities of plant nutrients but by the rapidity with which these nutrients are liberated in an available form. A knowledge of some of the more fundamental microbial processes is of prime importance for the proper application of certain principles relative to good soil-management practices. Several questions will serve as an outline for this study.

#### Questions

1. What are the most important beneficial effects of soil organisms?
2. What changes are brought about in the soil organic matter and its decomposition products?
3. Is the fixation of atmospheric nitrogen a microbiological process?
4. How does nitrification differ from nitrogen fixation?
5. By what process are nodules produced?
6. How may legumes be inoculated?
7. Activities of *Rhizobia* are influenced by what soil conditions?
8. How is fixed nitrogen utilized?
9. How much nitrogen is fixed through legume growth?
10. What soil microbial processes are detrimental?

**Beneficial Functions of Soil Microbes.** The beneficial effects of various microbes may be conveniently discussed under the following

three general topics: (1) changes brought about in the soil organic matter and its decomposition products, (2) fixation of atmospheric nitrogen, and (3) changes brought about in the soil mineral constituents.

**Changes in Organic Matter and Its Decomposition Products.** One of the chief activities of microorganisms is the mineralization of organic matter. By mineralization of organic matter is meant the breaking down of plant and animal organic materials into simple inorganic elements or compounds through the activities of the various organisms. Plant or animal residues, as such, cannot be used by higher plants, but through decomposition by microorganisms they become a valuable source of nutrients. All forms of soil organisms are concerned with the processes of organic matter decomposition, but the actions of bacteria, fungi, and actinomyces are the most important. These organisms bring about various transformations such as hydrolysis and oxidation by means of enzymes, which are usually called organic catalysts. The physical and chemical changes which organic materials undergo when changed to simple compounds by soil organisms are termed *decomposition*. The processes of decomposition vary greatly, however, with the nature of the organic substance and various other conditions, especially the degree of aeration.

When organic material is added to soils in the presence of sufficient oxygen (aerobic conditions), the non-nitrogenous compounds, such as starches, sugars, lignocellulose, organic acids, and oils, are decomposed by microorganisms and converted into various intermediate simpler compounds. But, regardless of the kind of intermediate products formed from the non-nitrogenous organic compounds, decomposition leads to the formation of a succession of simpler compounds. The carbon, hydrogen, and oxygen contained in the plant and animal materials are finally reunited to form carbon dioxide and water; the phosphorus is combined with oxygen to form phosphoric acid; and the sulfur with oxygen to form sulfuric acid. The mineral constituents of the original plant or animal material are left as a residue, mainly as salts of calcium, magnesium, potassium, and sodium. Examples of such salts are calcium phosphate, magnesium sulfate, potassium carbonate, and sodium chloride.

Decomposition in the absence of sufficient oxygen (anaerobic conditions) is brought about by entirely different kinds of organisms from those that function under aerobic conditions, and the chemical changes produced are not the same. The rate of decomposition is much slower, and the chemical changes are less complete. Although some carbon dioxide, water, and ammonia are formed from the complex organic

materials, most of the nitrogen, carbon, hydrogen, and oxygen are found in the form of more or less complex intermediate compounds, many of which are quite resistant to further decomposition. Under anaerobic conditions many compounds having offensive odors and a few poisonous compounds are produced. In the list of such compounds may be found hydrogen sulfide, phosphine, skatole, indole, and methane.

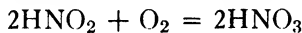
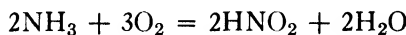
Owing to the great diversity of conditions affecting microbial activity and to differences in the nature of the organic materials, the processes of decomposition of organic matter in soils are of extreme complexity. When fresh organic substances are added to soils, the numerous bacteria, fungi, protozoa, and other organisms attack the various constituents, breaking them down into more simple compounds. During the processes of decomposition the chemical changes as a rule go through an orderly progression although many of the changes may occur simultaneously.

*Ammonification.* Ammonification may be defined as the formation of ammonia by soil organisms as a product of the decomposition of organic nitrogenous compounds. Ammonia is formed by a large number of soil organisms, including various fungi, actinomycetes, and aerobic and anaerobic bacteria. Thus the soil organisms responsible for ammonification are widely distributed and are found in abundance in most soils. The rate of ammonification and the quantity of ammonia produced are influenced markedly by soil conditions, nature of the organic material, and nature of the microbial population. The soil-management practices that especially favor ammonification are liming acid soils, draining wet soils, adding easily decomposable organic matter high in nitrogen, and proper tillage. The process of ammonification is of extreme importance in crop growing not only because certain plants have the ability to use ammonia compounds directly but also because ammonification is a necessary preliminary step to nitrification.

*Nitrification.* Another beneficial effect of soil organisms is the conversion of ammonia nitrogen to nitrate nitrogen—the process of nitrification. The fact that the nitrates, produced in the soil, are the principal sources of available nitrogen for practically all agricultural plants makes this a most important biological process. The conversion is carried on mainly by two specific types or groups of bacteria, of which there are several species, often referred to as the nitrifiers or nitrifying bacteria. Although seldom ever found in great abundance in any soil, fortunately they are widely distributed, being present in every cultivated soil. As would be expected, they are usually more numerous and especially more active in rich than in poor, unproductive soils.

The conversion biologically of ammonia into nitrate nitrogen is the result of two distinct processes. First, the ammonia is changed to nitrite nitrogen principally by the action of two groups of bacteria called *Nitrosomonas* and *Nitrosococcus*, and, secondly, the nitrite nitrogen is changed to nitrate nitrogen by the action of a group of bacteria called *Nitrobacter*. These nitrifying organisms use the energy liberated (as heat) in the nitrification processes for their growth and development, and the carbon for the synthesis of their cells is taken from the carbon dioxide of the atmosphere.

The reactions indicating the two steps in nitrification may be represented as follows:



These reactions show the change to be an oxidation process. It is probable that the  $\text{HNO}_2$  is neutralized by combination with a basic element such as calcium before being oxidized to the nitrate form. As a rule, in soils of humid regions, there is no appreciable accumulation of either ammonia or nitrites because they are rapidly oxidized. It is apparent that the process of nitrification does not increase the total supply of nitrogen in the soil; merely the form of nitrogen is changed.

In good agricultural soils nitrous acid and nitric acid as such probably do not exist, but they are combined with basic elements, forming salts. For example, nitrate nitrogen in soils usually is found as the nitrates of calcium, potassium, magnesium, and sodium, depending on the chemical condition of the soil.

The soil conditions under which active nitrification occurs are much more restricted than those for ammonification, because of the limited number of types of organisms concerned with nitrification and because the nitrifying organisms in general have similar growth characteristics. Consequently, the process is affected much more by changes in soil reaction, aeration, moisture, salt concentration, and other environmental conditions than is ammonification. Perhaps the most important factors which affect the activity of nitrifying bacteria in cultivated soils and which the farmer can more or less control are drainage, organic matter supply, fertilizers, and soil reaction. Drainage increases aeration, and, since nitrification is an oxidation process, the need for oxygen is obvious. Crops frequently suffer for want of available nitrogen in soils poorly drained owing to the inactivity of the nitrifying bacteria. The optimum reaction for the nitrifiers is pH 7.0 to 8.0; although some strains will

grow between the *pH* limits of 3.5 and 10.0, their activity is retarded in both strongly acid and highly alkaline soils.

*Sulfification.* Organic materials commonly added to soils contain compounds of sulfur, and, when the organic matter decomposes, the sulfur is released from its organic combination and eventually appears in an inorganic form, usually as  $H_2S$  and perhaps as some free sulfur. These inorganic products are oxidized by certain specific groups of bacteria into sulfurous acid ( $H_2SO_3$ ) and finally into sulfuric acid ( $H_2SO_4$ ). Some fungi are also believed capable of bringing about the latter reaction. The transformation of sulfur in organic and inorganic compounds to available sulfate forms by soil microbes is called sulfification. It is the same type of chemical change (oxidation) that takes place in nitrification. When sulfurous acid is formed, it usually combines with some of the soil mineral elements to form sulfites of calcium, magnesium, sodium, and potassium. They are then oxidized to sulfates. The importance of the sulfification process is realized when it is remembered that the higher plants take in most of their sulfur in the sulfate form.

*Assimilation and Synthesis of Organic Matter Decomposition Products.* If soil microbes have access to an abundance of food and energy materials, and if environmental conditions are favorable, they build up a considerable amount of organic matter of their own in the form of synthesized cell substance. In so doing they assimilate certain simple soluble products formed as a result of their own activities. In other words, large quantities of nitrogen, phosphorus, and other nutrient elements may be temporarily stored in the bodies of soil microorganisms in their processes of synthesizing cell substance. This assimilation of soluble plant nutrients may or may not be harmful, depending on whether or not it occurs during a time when they are needed by higher plants. The process would be highly desirable where there is danger of leaching before the nutrients can be used by crops. The addition of carbonaceous material, such as straw, to soil may greatly stimulate microbial growth and materially decrease leaching losses of soluble plant-food materials. It is obvious that conditions frequently may arise where soil microbes offer serious competition with higher plants for soluble nutrients. This is especially true of nitrate nitrogen.

Another important function of soil microbes, related to the assimilation and synthesis of organic matter decomposition products, is associated with the formation of humus, which is discussed in Chapter 10.

**Fixation of Atmospheric Nitrogen.** Nitrogen in its elemental form is a colorless, odorless gas and very inert. There is an inexhaustible

supply of it in the air, but it is in the free state and does not easily combine with other elements. It is estimated that there is the equivalent of 148,000 tons of nitrogen in the atmosphere for every acre of land area, the atmosphere being approximately 80 per cent nitrogen. Regardless of the fact that nitrogen is so inert in the free state, certain groups of soil organisms have the ability to take that element out of the air and utilize it in the building of their cells. The nitrogen of the air is thereby changed to a "fixed" form in which it can be of subsequent use to higher plants. This changing of atmospheric nitrogen into nitrogen compounds in the soil by microorganisms is known as *nitrogen fixation*. The process is accomplished mainly by two groups of bacteria, symbiotic and non-symbiotic, and there are several species of each.

*Non-Symbiotic Nitrogen Fixation.* There are certain groups of heterotrophic bacteria living in the soil independently of higher plants that have the ability to use atmospheric nitrogen in the synthesis of their body tissues. Since these bacteria do not grow in association (mutual relationship) with higher plants, they are termed non-symbiotic. These bacteria may be separated into two groups, on the basis of their oxygen requirements: (1) the aerobic group and (2) the anaerobic organisms. The former are known as *Azotobacter* and the latter as *Clostridium*.

*Azotobacter* are widely distributed in nature. They have been found in soils (of pH 6.0 or above) in practically every locality where examinations have been made. The greatest limiting factor affecting their distribution in soils appears to be the soil reaction. These organisms may exist in soils below pH 6.0, but as a rule they are not active, as far as nitrogen fixation is concerned, under such conditions. *Azotobacter* are favored by good aeration, abundant organic matter (particularly of a carbonaceous nature), the presence of ample available calcium and sufficient quantities of available nutrient elements, especially phosphorus, and proper moisture and temperature relations. So responsive are *Azotobacter* to phosphorus supply that they have been used as an agent in testing soils for the quantity of available phosphorus. The individual farmer, by applying lime and fertilizers and by returning as much of the crop residues as he can, may make the soil conditions more favorable for these nitrogen-fixing bacteria. Attempts have been made from time to time to increase the crop-producing power of soils by inoculating them with *Azotobacter*. For the most part, this practice has not been justified because, unless soil conditions are favorable, especially with respect to lime, the bacteria will not survive or function and, if soil conditions are

made favorable, the organisms will naturally become established without the aid of man.

The anaerobic bacteria, *Clostridium*, are much more acid-tolerant than members of the aerobic group and perhaps for that reason are more widespread. It is believed that these organisms can be found in every soil and that under suitable conditions they fix some nitrogen. It is not necessary that soils be waterlogged in order for anaerobic bacteria to function. A soil in good tilth may contain considerable areas within the granules favorable for the activities of these anaerobic nitrogen-fixing bacteria.

It is not to be inferred from this brief discussion of non-symbiotic nitrogen fixation in soils (frequently called azofication) that *Azotobacter* and *Clostridium* are the only soil organisms that have the ability to take nitrogen from the atmosphere and incorporate it in their bodies. It is probable that they are responsible for most of the non-symbiotic fixation; however, possibly several different groups of bacteria and perhaps fungi have that capacity. There is also definite evidence that certain species of blue-green algae have the ability to gather nitrogen from the atmosphere.

A question that naturally comes to mind is, how much nitrogen is fixed per acre per year by these non-symbiotic microorganisms under favorable field conditions? This question cannot be answered definitely because of the many difficulties encountered in making such a measurement under field conditions. However, in fallow soils increases in nitrogen have been secured which can be accounted for only through the activities of non-symbiotic organisms. These organisms can be isolated from soils, and they will fix nitrogen under laboratory conditions; it is therefore assumed that they will do likewise under field conditions, and thus they must be responsible for observed increases in soil nitrogen. Although accurate measurement of the quantity of nitrogen fixed cannot be made, the best estimates range from 10 pounds to 50 pounds per acre per year. The importance of this biological process to agriculture is realized when one considers that the amount so fixed, if an average fixation of 20 pounds per acre per year is assumed, is equivalent to the nitrogen in a 100-pound application of ammonium sulfate.

*Nitrification vs. Nitrogen Fixation.* Nitrogen fixation is fundamentally a different process from nitrification. Nitrogen fixation increases the soil's supply of nitrogen, whereas nitrification merely changes the compounds in which the nitrogen is held. The quantities of this element taken from the air and synthesized into proteins and other organic compounds by nitrogen-fixing bacteria are finally broken down by various

heterotrophic organisms to ammonia, which in turn undergoes nitrification. Thus, in general, nitrogen of the air goes through the processes of fixation, synthesis into proteins, decomposition, and ultimately nitrification before higher plants can use it. It should be mentioned, however, that certain soluble nitrogen compounds may be secreted from the bacterial cells during their active growth period and in turn may be utilized directly by higher plants. In other words, it may not always be necessary for the cells of nitrogen-fixing bacteria to decompose in order for higher plants to derive benefit from the nitrogen-fixation process.

*Symbiotic Nitrogen Fixation.* The most important bacteria, from the agricultural point of view, capable of utilizing the free nitrogen of the air are those that cause the formation of nodules on the roots of legumes. These organisms, when growing in the nodules of legume plants, derive their food, energy, and minerals from the legume, and in turn they supply the legume with some of its nitrogen. This growing together for a mutual benefit is called *symbiosis*, and hence the organisms are designated symbiotic nitrogen-fixing bacteria. The economic importance of symbiotic fixation is realized when it is said that the extensive growing of legumes is a major factor in the attainment and maintenance of a high level of agriculture. It has been estimated that nearly 2,000,000 tons of nitrogen are fixed annually by legume bacteria in the United States.

*Nodule Formation.* When these bacteria come in contact with the roots of the legume plant, some of them enter the single-celled root hairs. A rapid increase in the growth rate and in the numbers of bacteria then takes place because of the abundance of easily accessible food. These bacteria form an infection thread toward the base of the root hair that eventually penetrates the cortex of the root. This infection injures the legume plant and, in response to this stimulus, numerous plant cells in the meristematic tissue are produced in the immediate vicinity of the infection, eventually forming the nodule. The nodules, then, are essentially nothing more than masses of root tissue in which the bacteria live. The nodules are connected with the vascular system of the root; thus materials are transported to and from them. Each nodule may contain millions of bacteria, and the number of nodules that may develop on a single plant may vary from a few to a thousand or more.

Nodules are usually found in bunches, mainly on the younger roots; or in annual legumes they are frequently distributed about the tap root or the first-formed lateral roots.

The individual nodules may vary greatly in size and shape on different kinds of legumes. For example, the cultivated annual legumes generally have large sphere-like nodules, whereas those on the biennial and perennial legumes tend to be smaller, elongated, and in clusters.

Occasionally, nodule-like growths may be produced by nematodes or by crown-gall bacteria on the roots of legumes. These false nodules should not be confused with true nodules when determining the degree of infection by legume bacteria. Furthermore, certain non-leguminous plants frequently are found to possess nodule-like growths, which are

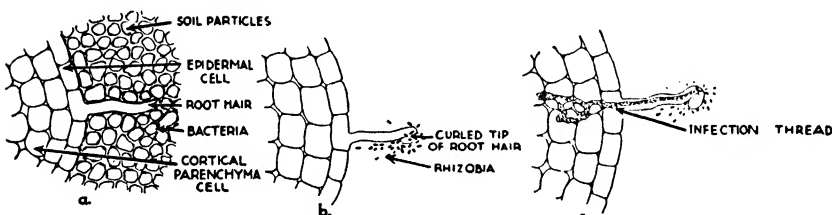


FIG. 47. Early stages in the formation of a nodule. (a) Response of the bacteria to a product of the host plant; organism moves toward root hair. (b) Curling of the root hair. (c) Early penetration of the infection thread. [Courtesy of P. W. Wilson and the University of Wisconsin Press, *The Biochemistry of Symbiotic Nitrogen Fixation*, 1940.]

produced by mycorrhiza, crown-gall organisms, and certain nematodes; but on examination they are not difficult to distinguish from the leguminous nodules.

*Inoculation for Legumes.* In order to obtain the full benefits from the growing of legumes, it is necessary that they possess true nodules. If active nodule-forming bacteria are not present in soils, they should be supplied—a process known as inoculation. A convenient and generally satisfactory method of inoculating a soil is to use pure cultures of legume bacteria, usually prepared in liquid and dust form or on agar. The liquid and dust cultures are ready for immediate use, but the agar cultures first must be shaken with water to obtain a suspension of the bacteria. The liquid or the suspension is applied to the seeds, which are allowed to dry in the shade before being sown.

No one particular kind of bacteria can be used to inoculate all legumes successfully. Different plant species as a rule possess their own particular bacteria, although sometimes one kind of organism may inoculate a half dozen or more plant species. Legume bacteria have been classed in the genus *Rhizobium*, of which there are several species and

strains. Some of the more common groups, each group representing a separate species of bacteria, follow:

1. Alfalfa group (*Rhizobium meliloti*), including bacteria that will produce nodules upon white sweet clover and yellow sweet clover as well as alfalfa.
2. Clover group (*R. trifolii*), which will produce nodules upon clovers such as red, alsike, crimson, mammoth, and white Dutch.
3. Pea group (*R. leguminosarum*), containing bacteria that produce nodules upon garden and field peas, sweet peas, vetches, etc.
4. Bean group (*R. phaseoli*), including bacteria capable of producing nodules upon garden, kidney, and navy beans and scarlet runner.
5. Soybean group (*R. japonicum*), those bacteria that produce nodules on soybeans.
6. Cowpea groups (*R. \_\_\_\_\_*), including bacteria producing nodules upon cowpeas, peanuts, lespedeza, velvet beans, and lima beans.
7. Several other small and less important groups.

When a legume crop is planted on a piece of land for the first time, it is essential that the soil be inoculated with the proper bacteria. The fact that red clover may have been grown on the land and produced an abundance of nodules is no indication that, for example, alfalfa bacteria will also be present. When there is doubt, the land should always be inoculated because the expense is small compared to the cost for seed and preparation of the seedbed. If a legume crop is grown without proper nodule production, the full benefit from that crop has not been realized. If there is no nodule production, the effect of legume crops on the soil would be similar to that of non-legumes, for the plants must get all their nitrogen from the soil. Good inoculation, when needed, results in an increase in yields and nitrogen content of the crop grown.

*Soil Conditions Affecting Rhizobia.* The question is frequently asked, how often should I inoculate my soil for the growing of certain legumes? That depends largely on soil conditions. Some of the more important factors which influence the growth and longevity of nodule bacteria are air supply, temperature, sunlight, reaction associated with the calcium content, supply of mineral elements, and the antagonistic action of other soil organisms. It can be said that unless killed by unfavorable soil conditions, *Rhizobia* may live and maintain their vitality in a soil for a number of years, possibly 10 to 20 years, even in the absence of the particular host legume. Generally, however, the legume bacteria in a soil seem to be rejuvenated by association with their host plant every

few years, or the effectiveness of the bacteria can be increased, as evidenced by increased nodule production and nitrogen fixation, by reinoculation. It is not to be inferred that there is always a correlation between numbers or quantities of nodules and the amount of nitrogen fixed, because certain strains apparently have the ability to produce nodules but do not gather nitrogen. Of the various factors affecting the longevity of legume bacteria in normal soils, their ability to produce nodules and their effectiveness in fixing nitrogen, calcium supply seems to be the most important. A good supply of active calcium not only in the soil but also in the plant is necessary for proper nodulation and active nitrogen fixation. In most soils available calcium seems to be more of a controlling factor in this respect than does pH.

*Utilization of Fixed Nitrogen.* The nitrogen gathered by legume bacteria may be utilized in three ways. It may be absorbed by the legume plant itself, it may be excreted from the nodule into the soil and be utilized by some crop growing in association with the legume, or when the legume crop is plowed down or dies the nitrogen may be released after the decomposition of the nodules.

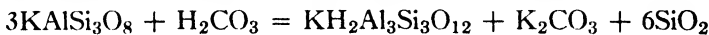
*Quantity of Nitrogen Fixed.* The quantity of nitrogen added to the soil through growth of legumes varies greatly according to many conditions, such as the kind of legume, the nature of the soil, the effectiveness of the bacteria present, and seasonal conditions. In general, it can be said that, the poorer the soil in available nitrogen and the richer in lime and available phosphorus and potash, the greater will be the gain in nitrogen by growing legumes. It appears that the intimate relations existing between nodule bacteria and their host plants are determined mainly by the carbohydrate supply in the host plants. Any environmental condition affecting the production of carbohydrates in the plant would automatically affect the quantity of nitrogen fixed by good strains of legume bacteria. Well-inoculated legumes growing under favorable conditions will usually gather 50 to 100 pounds of nitrogen per acre per year. With good crops of such legumes as alfalfa and sweet clover the quantities fixed may be even greater than 100 pounds per acre per year.

In actual farm practice the amount of nitrogen added to a soil by legume bacteria is determined by the methods of disposing of the legume crop. If the crop is turned under as a green manure, the total quantity of nitrogen taken from the air is added. If the crop is cut for hay and sold off the farm, little or no gain is realized; with some legumes there may even be a net loss of nitrogen. And, if the crop is cut for hay and fed on the farm, about one-half the nitrogen that was taken from the

air by the legume bacteria can be returned to the soil if special care is exercised in handling the manure to prevent loss. It is generally assumed (although not necessarily true for all legumes) that the amount of nitrogen in the roots and stubble equals the amount of nitrogen taken from the soil; this would mean that the quantity of nitrogen removed by the harvested crop is equal to the nitrogen obtained from the air.

**Changes in Inorganic Soil Constituents by Microorganisms.** It becomes evident that in the decomposition of organic matter and in the synthesis of microbial cells there are numerous reactions involving the solubility of various soil mineral elements. Among those mineral elements which are affected directly or indirectly by the action of microorganisms, the following should be mentioned: phosphorus, sulfur, calcium, potassium, magnesium, iron, manganese, aluminum, and silicon. Sometimes soluble compounds of the elements are formed, whereas at other times insoluble compounds result.

The most common acids produced in the soil as a result of the action of microorganisms are nitrous, nitric, phosphoric, sulfurous, sulfuric, carbonic and numerous other organic acids. The following changes result from the action of these acids: (1) insoluble phosphates are changed to more soluble compounds; (2) calcium, magnesium, and other slightly soluble carbonates are changed to more soluble forms; and (3) insoluble potassium compounds are also changed to more soluble combinations. The reactions for some of the above changes may occur according to the equations



Practically all mineral elements related to the growth of higher plants either as nutrients or "stimulants" are subject to the action, in one way or another, of microorganisms in the soil.

These few statements show that extensive and far-reaching chemical changes are produced in the inorganic soil constituents through the various activities of soil microorganisms. They are the primary agents which convert the elements of the insoluble mineral and organic compounds into soluble forms available as plant nutrients.

**Soil Microbial Processes of a Detrimental Nature.** For the most part microbial processes are highly beneficial, but certain of the detrimental effects should not go unmentioned. The harmful activities of

soil organisms may be grouped as follows: (1) the production of plant and animal disease, (2) denitrification, (3) competition with higher plants for available nutrients, and (4) the production of toxic compounds.

*Production of Disease.* The soil frequently contains a rather large number of organisms that cause diseases either in plants or animals. Some of these organisms live in the soil only temporarily, and others use it as a permanent habitat. The soil may harbor organisms that cause such bacterial diseases as wilt of tomatoes and potatoes, soft rots of a number of vegetables, leaf spots, and galls. Some of the most destructive parasites are the disease-causing fungi such as those that cause damping-off of seedlings, cabbage yellows, mildews, blights, certain rusts, wilt diseases, scab, dry rot of potatoes, and many others. Certain species of *Actinomyces* may cause diseases like scab in potatoes and sugar beets and pox in sweet potatoes. Nematodes of a parasitic nature commonly infest soils and cause injury to many plants, especially to the roots. Abnormalities or swellings on the roots, usually referred to as root knot or root gall, are produced.

So-called "soil sickness," frequently encountered in greenhouses, is often due to the presence of parasitic soil organisms of one kind or another. The ability of soil-borne microbes to infect a plant and cause disease is often a function of the temperature, moisture, nutrient, or acidity relationships. Partial or complete control may be exercised by regulating or modifying the soil conditions, the theory of control then being that the parasitic organisms function through a narrower range than that of the higher plants which they infect.

*Denitrification.* Denitrification may be considered the reverse of nitrification. It is a reduction process in which oxygen is taken away from nitrates to form, first, nitrites, then either ammonia or gaseous nitrogen. In the ammonia or gaseous nitrogen form, the nitrogen may escape into the atmosphere. This destructive or unfavorable process takes place in the absence of oxygen and in the presence of an abundance of soluble organic matter, which serves as a good source of energy for the organisms concerned. The fact that denitrification takes place in the absence of air explains the decomposition of nitrates in the lower layers of soils as well as in surface soils that are deficient in oxygen. Furthermore, that is why heavy, compact soils tend to suffer greater loss from denitrification than do the more open, sandy soils. Any soil-management practice that promotes the circulation of air in soils lessens the danger of denitrification. In ordinary cultivated soils denitrification is of little economic importance as a rule,

although, if a cropped field becomes flooded for a considerable length of time, a considerable loss in nitrogen may result from denitrification.

*Microbial Competition with Higher Plants for Available Nutrients.*

Attention has already been called (p. 216) to the beneficial effect of the assimilation of nutrient elements by microorganisms in decreasing leaching losses, but a pertinent question now arises: Do microorganisms seriously compete with higher plants for nutrients? The soil organisms require nitrogen, phosphorus, potassium, calcium, and other elements, as do higher plants, and obviously even a slight increase in the microbial population might appreciably reduce the available nutrients of the soil to a point at which the growth of higher plants would be impaired. Aside from carbon, hydrogen, and oxygen, nitrogen and phosphorus are the elements usually consumed in largest quantities by microorganisms and the elements for which the keenest competition between microbes and higher plants exists. Of the two elements, nitrogen is more vigorously competed for.

If straw or any easily decomposable carbonaceous material is added to a soil containing a liberal supply of nitrates, biological activities are greatly accelerated. The soil organisms, having an abundant supply of food, will multiply rapidly and will need a large supply of nitrogen for the synthesis of their protoplasm. Since the carbonaceous food material will not supply the necessary amounts of nitrogen, the organisms will utilize the soluble nitrates of the soil, thereby decreasing the supply available to higher plants. Conditions may thus exist whereby higher plants suffer seriously for available nutrients, owing to competition of the soil microorganisms. For this reason, crops following immediately the turning under of straw or similar material may suffer for the want of available plant nutrients. The remedy then would be the liberal application of a fertilizer supplying these nutrients. Soluble nutrients taken up by microorganisms are "tied up" in an unavailable form only temporarily. When the organisms die and their bodies decompose, the plant nutrients are again released in an available form.

*The Production of Toxic Substances.* Substances toxic to plant growth as well as to certain microorganisms accumulate, under anaerobic conditions, in some soils as a result of the incomplete decomposition of organic matter. For example, methane, hydrogen sulfide, phosphine, skatole, indole, and numerous organic acids are toxic compounds produced, but they are generally considered merely products of improper soil conditions and when these soil conditions are corrected the toxic materials disappear. If good drainage and tillage are provided, together

with the proper use of fertilizers and lime, these toxins will not be a problem in soils.

### THE ROLE OF MICROORGANISMS IN THE DEVELOPMENT OF SOILS

Without living organisms there could be no natural soils, and hence no one will question the contribution of biological processes in soil formation. The earliest students of soils were fundamentally students of geology and chemistry, and consequently they did not pay much attention to the roles played by organisms in rock weathering and soil development. Recently more consideration has been given to the part played by microorganisms, and it is suggested that, possibly, biological processes are of equal, if not of more, importance in soil formation than are the non-biological processes. It is obvious that the biological and non-biological forces act together. Answers to the questions below should be of interest to the student of soils.

#### Questions

1. How do the microorganisms contribute to the decomposition of rocks and minerals?
2. Is decomposition of soil organic matter related to soil development?
3. Do microorganisms influence soil-leaching losses?
4. Is the movement of soil material within the profile affected by microorganisms?
5. Do organisms aid in the development of a good soil structure?
6. In what ways may the animal population affect soil development?
7. What is the main driving force of the biological factors of soil formation?

**Decomposition of Rocks and Minerals.** It has already been mentioned that microbes are capable of bringing about extensive changes in a number of minerals and compounds of mineral elements. They assimilate such elements as phosphorus, calcium, magnesium, iron, and potassium from slightly soluble mineral compounds. They produce various organic and inorganic acids which exert a solvent action on most minerals. Soil organisms such as the algae and lichens have a corroding effect on rocks and minerals because of the production of acids. Other organisms increase the solubility of feldspars and other silicates, and many of them are active in promoting the mechanical weathering of minerals.

**Organic Matter Decomposition.** The amount and nature of the organic matter and its decomposition are of major concern in the formation of a soil, and the control of organic matter decomposition is exercised to a large extent by the microorganisms. In the cycle of mineral elements which are taken up by plants from the lower soil horizons and then returned to the surface horizons, there exists a close relationship to the microbial population. Decomposition and mineralization, as has been pointed out, are important functions of the soil organisms whose activities are largely controlled by climatic conditions. In cool, humid climates biological activity is slowed down and decomposition of organic matter is slight; these conditions permit considerable accumulation which in extreme cases may result in the formation of peat or tundra soils. In warm, humid climates, on the other hand, rapid decomposition does not permit a great accumulation of organic matter in well-drained soils. The role of humus in soil formation is discussed elsewhere (pp. 33, 42).

**Leaching of Soil Materials.** The function of microbes in affecting leaching losses is closely associated with the decomposition of minerals and organic matter. Leaching losses are greatly enhanced by the activities of microorganisms, owing to their effect in increasing the solubility of both mineral and organic constituents. The leaching process is of major importance in determining the nature of the soil profile that is developed. It is to be pointed out in this connection, however, that certain quantities of the nutrient elements which are made soluble are assimilated by the microorganisms and are not immediately lost by leaching.

**Movement of Soil Material.** Microorganisms play a part in the translocation of soil material through the production of substances which exert a solvent effect on certain soil constituents. The dissolving effect of organic and inorganic acids on more or less insoluble compounds is obviously a contributing factor in the translocation of some products by soil-moisture movements. Furthermore, some of the mineral elements that are released combine with the organic colloidal constituents and move in the colloidal state. Products of microbial activity may appreciably affect the dispersion of soil colloids and thus influence their movement in the soil profile. Also, modification of the soil reaction not only will affect the solubility of certain soil constituents but also will have a pronounced influence on soil structure, both of which actions are concerned in the movement of soil material.

**Structure of Soils.** In the development of soil structure, micro-organisms play an important but indirect part. They are responsible mainly for the formation of organic colloidal material which promotes the formation of a granular structure which is so desirable in soils. Some of the animals influence soil structure directly. For example, soil which is passed through earthworms is excreted in a highly granular condition.

**Effects of the Animal Population.** The work of the various animal species as soil formers is primarily mechanical. Ants, worms, termites, and rodents, for example, are instrumental in mixing the soil, thereby distributing the microbial population and incorporating organic matter with the mineral material. This action is of especial significance in timbered soils. Burrows made by these organisms also increase aeration and drainage. Chemical effects come about largely through the decomposition and mineralization of substances composing the bodies. Some chemical action may also take place when soil materials pass through the digestive system or when some chemical substance, such as an organic acid, is produced by the animal.

**General Remarks.** In relation to soil formation and development, enough has been said to indicate definitely that microbes through their diverse activities play parts of major importance. Sometimes they aid the purely chemical, physical, and mechanical changes; and sometimes they are the primary agents in the transformation processes; and at other times they play a prominent role, as in the formation of forest soils and of peat and muck soils. The main driving force behind the various biological factors is the *climate*.

#### RELATIONSHIPS BETWEEN HIGHER PLANTS AND SOIL MICRO-ORGANISMS AND AMONG SOIL MICROORGANISMS THEMSELVES

Neither microorganisms nor higher plants can develop normally for any length of time under natural conditions in the absence of the other. Even a very superficial examination of the relationships of microorganisms to higher plants brings to our minds a large number of interrelated factors, the most important of which are suggested by the questions below.

#### Questions

1. Does an increase in microbial activities accompany the growth of higher plants?

2. What is meant by the antibiotic effects of microorganisms?
3. Are microorganisms concerned with the production of growth-regulating substances in soils?

**Effect of Higher Plants on Microbial Activities.** The growth of higher plants in soils increases many microbial activities, and it is noteworthy that their activities become especially intense in the vicinity of the roots. The enhancement in microbial action begins when the higher plants are young and continues throughout the active growth period of the plants. This modification in the microbial flora is due in part to physical changes produced in the soil environment by the plant roots, but the most pronounced influence is believed to be exerted by the organic substances, arising from the roots themselves, which serve the heterotrophic organisms as a source of food and energy. Evidently it is the soil of the root zone that plays the most important part in the nutrition of higher plants, and it is here that microorganisms exert their most pronounced effects. It is in the root zone (*rhizosphere*) that microbes are most active in increasing the availability of nutrients for plants, and it is here also that they may exert their most injurious effects under certain conditions.

**Antibiotic Effects.** Only a relatively few of the bacteria and other microorganisms that cause diseases in animals and man survive long in the soil. If these disease-producing organisms are introduced into the soil, they are inhibited or killed by antagonistic organisms which produce active chemical substances known as antibiotics.

The purifying effect of the soil organisms has been recognized for many years, but it is only recently that a systematic attempt has been made to use them to control infections in man and animals. To date, only comparatively few groups of soil organisms have been examined for their antibiotic properties.

Since the work of Pasteur it has been established that the great majority of bacteria causing the common infections rapidly disappears from the soil. The rate of destruction of these pathogenic organisms in the soil depends on such factors as the moisture content of the soil, the reaction of the soil, and the nature and abundance of its microbiological population. It has been found that in most soils a large proportion of disease-producing bacteria die out within 10 days under conditions favorable to the development of the antagonistic organisms. It should be mentioned, however, that certain pathogenic spore-forming bacteria may persist more or less indefinitely in soils, but these organisms are relatively unimportant in that infections caused by such organisms are readily subject to control.

The experimental evidence indicates that the destruction of pathogenic organisms in soils by other microorganisms is due to the production of specific antibiotic agents. There appears to be little evidence that would lead one to believe that the antagonistic substances exist as such in the soil; rather, the antagonistic organism produces substances which are responsible for the destruction of the pathogenic organisms.

The great importance of soil microorganisms in the destruction of disease-producing bacteria has thus become definitely established and is quite generally recognized. In this connection it is necessary to mention only three important antibiotics, penicillin, streptomycin, and tyrothrinacin, produced by soil microorganisms. Penicillin is produced by a large number of fungi belonging to the genera *Aspergillus* and *Penicillium* which are widely distributed in the soil. Streptomycin is produced by certain strains of an organism belonging to the actinomycetes, and tyrothrinacin is produced by a group of aerobic spore-forming bacteria. In addition to tyrothrinacin, a number of other antibiotics have been isolated from soil bacteria. Although these organisms may be found in other substrates, the soil is considered their natural habitat.

The action of antibiotics against bacteria and other microorganisms is selective in nature. The antibiotics act upon some bacteria and not at all, or to only a limited extent, upon other bacteria. Some are effective against fungi, and others are not.

The antagonistic effects among the soil organisms may be very common, although such effects are extremely difficult to isolate and demonstrate. It has frequently been observed that plants affected by disease are greatly influenced by soil conditions. This effect is perhaps reflected through the general activity of the microbial population. For example, a certain disease may be suppressed by the addition of fertilizers, manures, green manures, or crop residues, which increase the activities of the organisms other than those responsible for the disease.

It has been demonstrated that certain bacteria may be antagonistic to some species of actinomycetes causing potato scab and that some fungi and bacteria suppress the activities of organisms responsible for root rot of wheat. These effects are believed to be due, at least in part, to antibiotic agents.

**Growth-Regulating Substances.** During the past few years considerable attention has been given to the effect of vitamins, particularly vitamin B<sub>1</sub> (thiamine chloride), on plant growth. Favorable effects from the application of some of these growth-regulating substances to the soils have been observed, in certain instances, but differences of opinion exist concerning the real nature of these phenomena or their

importance in agricultural practice. It is established, however, that vitamin B<sub>1</sub> is essential for plant growth; but, because it is produced in the decomposition of organic matter, synthesized by certain micro-organisms and higher plants, it is not likely to be a limiting factor in plant growth under favorable soil and environmental conditions.

The beneficial effects of organic matter in some soils have been ascribed to growth-regulating substances of a vitamin-like nature produced by microorganisms in the decomposition of the organic materials. As is generally known, the power to synthesize vitamin B<sub>1</sub> is not limited to green plants but is also possessed by certain fungi and other micro-organisms. The importance of vitamin synthesis by microorganisms in the digestive tract of animals and man is well recognized. Perhaps the members of the vitamin B complex are the principal vitamins which are synthesized by microorganisms, although there is considerable evidence that other vitamins may be synthesized also.

# 10

## Soil Organic Matter

In the study of soils centuries ago, it was observed that their capacity to produce crops was more or less directly related to the amount of organic matter which they contained. Likewise, the farmer of today, in designating soils which he considers highly fertile, usually selects the dark-colored ones. He makes this choice because experience has taught him that such soils are usually more productive than the light-colored ones, and hence in selecting dark-colored soils the farmer is involuntarily paying tribute to the value of organic matter in soils.

In general, it can be said that organic matter exerts a controlling influence on soil properties, including productivity, and without it the surface layer of the earth could hardly be correctly designated as soil. The organic matter content of soil is one of our most important and also easily exhausted resources. This subject may be studied under the following headings.

### Objectives

- A. Sources and kinds of soil organic matter.
- B. Decomposition of organic matter and humus formation.
- C. Effects of organic matter on soil characteristics and productivity.
- D. Accumulation and loss of soil organic matter.

### SOURCES AND KINDS OF SOIL ORGANIC MATTER

The roots, leaves, and stems of plants in various stages of decay constitute by far the largest part of the organic material of the soil. In farm lands much of this comes from the root system of crops and the mature stems and leaves after the seed has been harvested. The practice is increasing, however, of planting crops to protect the soil from erosion and to be plowed under while still green. Farm manures also provide the opportunity of returning large quantities of plant material to the soil. These materials are worked into the soil to a considerable extent by

plowing or other tillage practices. In timber and grazing land the plant leaves and stems are deposited on the soil surface and become incorporated into the soil to a limited extent by natural processes. In these soils the nature of the root system and whether the plants are grasses or trees and shrubs determine largely the amount of organic material within the soil.

In addition to the remains of higher plants, the bodies of small organisms such as worms, insects, molds, fungi, and bacteria contribute appreciable quantities of organic matter to many soils.

### Questions

1. What kinds of crop residues are returned to soils?
2. Why are green crops desirable sources of organic matter?
3. Does manure supply much organic matter?
4. Are peat and muck good sources of organic matter?
5. Do fertilizers supply much organic material?
6. What is meant by artificial manure?

**Crop Residues.** Leaves, stubble, roots, and mature plant materials such as straw and the stalks of corn, cotton, grain, sorghums, and similar crops are included in this group. They decompose rather slowly, and little nitrogen or other plant-nutrient elements are released until after a considerable period has elapsed. During the early stages of decay of these materials a shortage of nitrogen for a crop growing on the land may develop because these residues do not contain enough nitrogen to supply the needs of the decay organisms. This condition can be remedied by application of a fertilizer containing considerable nitrogen, by applying manure, or by working the residues into the soil well in advance of the time a crop is planted.

**Green-Manure Crops.** These include the various plants grown for cover and green-manuring purposes. They aid in conserving soil organic matter by virtue of the fact that (1) they decrease erosion losses; (2) they conserve plant-food elements by decreasing leaching losses; (3) when turned under, they increase the supply of soil organic matter; and (4) with leguminous crops, the total supply of nitrogen is increased, provided that the legumes are well inoculated with effective strains of bacteria. The ability of leguminous crops to increase the nitrogen content of soils is one of the primary reasons why they are considered especially valuable as sources of organic matter. The difference between legumes and non-legumes is not only that they vary in nitrogen-gathering ability but also that legumes contain more nitrogen in proportion to their carbon content.

Green plants are an excellent source of active organic matter; that is, they decompose rapidly. This active decomposition is accompanied by a rapid release in available forms of nitrogen and of the mineral elements contained in them. The subject of green-manuring is discussed in considerable detail in Chapter 11.

**Farm Manures.** The value of farm manures for increasing productivity is well recognized. They are ideal sources of soil organic matter. A large portion of the nitrogen and an appreciable part of the mineral nutrients contained in manure are liberated rapidly in forms available for plant use, and a rather large amount of residual organic matter (humus) is formed. On the average, a ton of farm manure contains about 500 pounds of dry matter and around 25 pounds of plant nutrients (10 pounds N, 5 pounds  $P_2O_5$ , and 10 pounds  $K_2O$ ).

**Peat, Muck, and Forest Litter.** These materials (with the exception of fresh forest litter and raw peat) have already undergone extensive decomposition. These materials must be considered only as sources of humus and not as fertilizers. They have a very low content of phosphorus and potassium, and the nitrogen is only slowly available. From the general farming point of view these materials have limited use. They are frequently used in the preparation of lawns, golf courses, and greenhouse soils.

**Organic Fertilizers.** This group of materials includes all fertilizers of plant or animal origin, such as dried blood, tankages, cottonseed meal, and sewage sludge. Generally speaking, they decompose rapidly and leave a rather small quantity of organic matter. The nitrogen in most of these materials is made available rather rapidly. Only in special cases can these materials be considered important sources of organic matter because the quantity usually added to soils is so small.

**Artificial Manures.** Composts of plant residues are excellent sources of organic matter if they have been well prepared and if sufficient nitrogen, phosphorus, and lime have been used to bring about rapid decomposition. When compost materials are added to soils, they produce or leave a considerable amount of organic matter in the soil and some of the nutrients they contain are readily liberated in an available form.

Artificial manures are produced by composting plant residues, such as leaves, straw, corn stover, grass, and weeds, under favorable conditions of moisture, temperature, and aeration. These residues are low in nitrogen, phosphorus, and lime, which are necessary for rapid decomposition of the organic materials by microorganisms. These three elements (N, P, and Ca), in readily available substances, must

be added to the compost. Several formulas have been suggested for chemical mixtures that are satisfactory for composting purposes. The following formula has been used with considerable success:

| (By Weight)               |          |
|---------------------------|----------|
| Ammonium sulfate          | 45 parts |
| Superphosphate (20%)      | 15 parts |
| Limestone (finely ground) | 40 parts |

This mixture is used at the rate of 150 pounds per ton of dry straw. A ready-mixed fertilizer such as a 10-6-4, in addition to lime, can be substituted for the chemical mixture given above. From 100 to 125 pounds of the 10-6-4 fertilizer and 50 or 60 pounds of finely ground limestone should be used per ton of straw. As the compost pile is built up, the fertilizer can be mixed with each layer of organic material. The time required to produce a good grade of manure is determined mainly by the conditions of temperature, moisture, and aeration in the compost pile.

#### DECOMPOSITION OF ORGANIC MATTER AND HUMUS FORMATION

If the crude organic materials added to soils in plant and animal remains did not decompose, they would be of little value in crop production or soil improvement. The plant nutrients in these materials are converted into a usable form (become available) only through decay processes. Likewise, the tissue of higher plants and animals, in its natural structure, contributes little to the physical or chemical improvement of soils. These structures must be broken down and the chemical make-up changed before they serve to improve soil tilth and aid in desirable chemical processes in the soil. A number of questions may serve as guides in the consideration of the decomposition of organic materials in the soil.

#### Questions

1. What is the chemical nature of organic matter added to soils?
2. What factors determine the rate of organic matter decomposition?
3. Which soil organisms are active in the decay process?
4. How is humus formed, and what is it?
5. What are the chief characteristics and properties of humus?
6. In what way are the nitrogen compounds in organic matter changed during organic decay?

7. Are the mineral constituents changed during organic decay?
8. Do organic compounds serve directly as plant food?

**Composition of Organic Materials Added to Soils.** Numerous substances synthesized by plants are added to soils in residues and green plants. Animal tissue and more or less decomposed plant tissue are supplied in manures and in industrial products such as dried blood, meat scrap, bonemeal, and the hulls and meals from various seeds as cotton, flax, and soybean. Large numbers of small animals find the soil either a temporary or permanent habitat, and a considerable quantity of microbial cell substance is continuously being added to the soil. Thus, we find in the soil any chemical compounds contained in higher plants, microorganisms, or animals. Needless to say, the composition of fresh organic materials added to soils is of extreme variability; this would be true even if the plant materials alone were considered. A knowledge of the chemical composition of plant residues undergoing decomposition is important in aiding us to understand the nature and rapidity of the liberation of the nutrient elements in forms available for plant growth and of the formation and nature of the residual organic matter.

Green plant tissue contains on the average about 75 per cent water. The dry matter, making up about 25 per cent of the plant, is composed mostly of carbon, hydrogen, and oxygen. Over 90 per cent of the dry-matter weight is furnished by these three elements, the remainder consists of nitrogen, sulfur, calcium, phosphorus, potassium, and other elements. Although the elements other than carbon, hydrogen, and oxygen make up only a small portion of the plant, they play an important role in plant nutrition as the plant residues are decomposed in the soil.

As a matter of convenience, the chemical compounds may be grouped into three general classes: (1) nitrogen-free organic compounds, (2) nitrogen-containing compounds, and (3) inorganic or mineral constituents. The compounds of the first group are composed of carbon, hydrogen, and oxygen and include the carbohydrates such as sugars and starches, the celluloses, lignocellulose and lignin, fats or oils and waxes, and organic acids. The nitrogen-containing compounds are mostly proteins, and in addition to carbon, hydrogen, oxygen, and nitrogen they may contain sulfur, phosphorus, iron, and other mineral elements. The inorganic or mineral constituents consist primarily of compounds of phosphorus, calcium, magnesium, potassium, silicon, sulfur, aluminum, iron, and manganese. More than 60 elements have been detected in the chemical analyses of plants. Some of the mineral ele-

ments may exist in the plant as combinations with organic compounds; some may occur as organic salts and others as inorganic salts.

**Factors Affecting the Rate of Organic Matter Decomposition.** Since the decomposition of organic matter in the soil is a biochemical process, any factor that affects the activities of the soil organisms will necessarily affect the rate of organic matter decay. Several factors which have a bearing on the rate of organic matter decomposition may be placed in the three following groups: (1) the nature of the plant material (including such points as the kind of plant, age of plant, and chemical composition); (2) soil (including aeration, temperature, moisture, acidity, and fertility level); and (3) climatic factors (the effects of moisture and temperature are particularly influential).

In general, the younger the plant the more rapid will be its rate of decomposition when incorporated into the soil. This is largely due to the higher content of water-soluble constituents, a higher nitrogen content, a more narrow carbon-nitrogen ratio, and a smaller percentage of lignin and other constituents resistant to decomposition. Of the various plant and animal residues used to increase the organic content of the soil, as a rule those having the higher nitrogen content decompose most rapidly.

The influence of several soil factors on the rate of microbiological activities and on the accompanying rate of organic matter decomposition is discussed in considerable detail elsewhere (p. 209). Soil conditions which are favorable for the growth of the common crop plants are also favorable for organic matter decomposition. One of the most influential factors in this respect is the degree of soil aeration. Since organic matter decay is largely a burning or oxidation process, the greater the degree of aeration the greater, in general, will be the rate of decay. Soil moisture and especially soil temperature are controlled largely by climatic influences. With other conditions favorable, the rate of soil organic matter decay increases with increasing soil moisture to a point where aeration is inadequate. The rate of decay increases with increasing temperature to a certain limit, if all other conditions are assumed to be constant and favorable.

**Organisms Involved in the Decomposition of Soil Organic Matter.** When plant and animal residues are added to the soil under favorable conditions, they are subject to various changes involving chemical agencies and the activities of the various groups of soil organisms. The animal population aids in the breaking down of plant residues; it brings about a reduction in quantity and a change in the chemical composition of the residues. Furthermore, it causes a mechanical mix-

ing of the organic with the inorganic soil constituents. It is the fungi, bacteria, actinomyces, and protozoa, however, which usually accomplish the most pronounced changes in the organic residues. All the organisms active in the soil are associated either directly or indirectly, with the decomposition of organic matter, for it is on this material that most of them depend for food and energy.

The organic residues are not decomposed as a whole, but the chemical constituents may be attacked independently of one another. A particular group of organisms is usually associated with the active decomposition of one special group of compounds; some organisms perform definite or specific activities. Attack by soil organisms on organic materials seldom results directly in complete decomposition; various intermediate products usually are formed. Other members of the soil population which have the ability decompose the intermediate products. Sooner or later the original material loses its identity and is transformed to a variety of end products and *synthesized substances*. Soil organisms differ greatly in their food requirements; in other words, what may be a source of readily available food for one group may not be available to others. This makes possible the decomposition of whatever organic material may be added to or formed in the soil.

For additional information on the soil organisms involved in the decomposition of soil organic matter, see Chapter 9.

**Humus Formation.** Humus denotes soil organic matter which has undergone extensive decomposition. It is not a homogeneous compound; it has no definite chemical composition. It is a dark, heterogeneous mass consisting of the residues of plant and animal materials together with the synthesized cell substances of soil organisms. Humus is not static but dynamic in soils; it is continually undergoing change.

It has already been pointed out that, during the decomposition of plant and animal residues in soils, some organic constituents are more readily attacked than others and that some are extremely resistant to decomposition. The starches, sugars, proteins, and amino acids are rapidly attacked by a great variety of organisms and, associated with these changes, is a considerable synthesis of microbial cell substance. The celluloses and especially the hemicelluloses are decomposed rapidly by a rather large variety of microorganisms. Of the various plant constituents that exist in considerable quantity, the lignins are particularly resistant to decomposition under anaerobic conditions, but under aerobic conditions they undergo some change although not so extensively as the celluloses and hemicelluloses.

Thus, in the formation of humus from plant residues there is a rapid

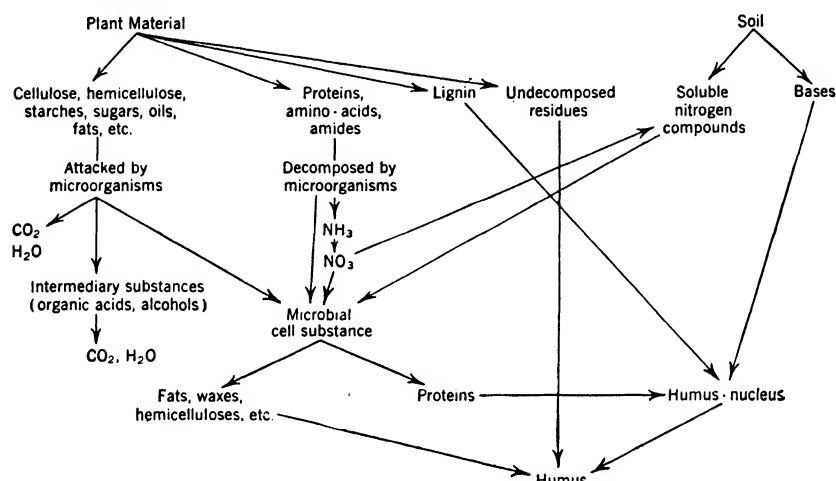
reduction of the water-soluble constituents, of the celluloses, and of the hemicelluloses; a relative increase in the percentage of lignin and lignin complexes; and an increase in the protein content unless the plant residues are unusually high in protein. The new protein is believed to be formed for the most part through the synthesizing activities of the microorganisms. The C:N ratio in fresh plant materials that are commonly added to soils is usually from about 80:1 in mature straw to 12 to 20:1 in leguminous green-manure crops (Table 28, p. 258). In any case, the ratio will narrow or drop to about 10:1 in a relatively short time after these materials have been incorporated into soils. In other words, the C:N ratio of humus in mineral soils is roughly 10:1. This means that for each pound of nitrogen in the soil there exist about 10 pounds of carbon.

The increase in the relative amount of nitrogen (as shown by the more narrow C:N ratio) in the organic residues after decomposition can only be explained by the fact that the nitrogenous complexes are rendered resistant to further rapid decomposition. This indicates, therefore, that either the organic nitrogenous complexes in humus are not of a true protein nature or they are not in a free state; otherwise they would decompose readily. This slow rate of decomposition of humus is obviously of considerable practical importance. It offers a means whereby nitrogen can be stored in the soil and released gradually.

Humus can be considered to consist largely of two chemical complexes—lignin and protein. Perhaps these two groups of substances combine, forming a lignin-protein complex. Lignin makes up 40 to 45 per cent of the total humus, 30 to 35 per cent of the humus consists of protein, and the remainder is composed largely of fats, waxes, and other residual materials. The lignin-protein combination renders the protein resistant to microbial attack. It is thus observed that humus has a high protein content; yet the nitrogen is only slowly available for the growth of higher plants. The lignin in humus originates mostly from plant residues with perhaps certain chemical modifications, whereas the proteins in humus are synthesized mainly through the activities of microorganisms. This combination of lignins and proteins has been called a ligno-proteinate complex. It is seen from the foregoing discussion that nitrogen is a key element in the formation of humus, and, because the crude plant materials applied to soil are relatively low in nitrogen, the quantity of humus produced from them is limited. The importance of lignin in the humification process must not be lost sight of, however, for unless lignin is present to combine with the nitrogen humus formation will not take place.

Although humus is considered organic, it probably contains various inorganic elements which are an integral part of the complex. Phosphorus, sulfur, calcium, magnesium, potassium, iron, aluminum, and possibly other elements may be bound chemically with the humus. In acid soils the humus complex is likely to contain large quantities of hydrogen, iron, and aluminum; whereas in soils close to the neutral point the humus usually is nearly saturated with calcium and mag-

**Chart 7. Schematic Representation of the Mechanism of the Formation of Humus in the Decomposition of Plant Residues in Soil**



[After Waksman, from *Humus*, by permission of William and Wilkins Co.]

nesium, and in strongly alkaline soils it may contain considerable sodium.

Since the lignin and protein account for 70 to 80 per cent of the total humus, the formation of a lignin-protein complex suggests evidence to explain the more or less constant C:N ratio which exists in mineral soils. These complexes, however, are not absolutely resistant to decomposition; therefore the nitrogen content of humus may not always be constant. Furthermore, the chemical nature of humus is determined to a certain extent by the chemical composition of the plant residues and by soil and climatic factors.

A general schematic summary of the processes leading to humus formation is presented in Chart 7. The material commonly referred to as humus includes the mass of plant residues undergoing decomposition, together with the synthesized cell substance and certain in-

termediary and end products. It is constantly changing in composition. It is better, therefore, to speak of humus not as a single group of substances but rather as a state of matter, which is different under varying conditions of formation.

**Characteristics and Properties of Humus.** In the preceding section, in the discussion of the formation of humus in mineral soils, it was pointed out that some plant constituents become completely decomposed, others are modified to a greater or lesser extent, and others are only slightly attacked, and that these processes are accompanied by the synthesis of new complexes (fats, carbohydrates, and proteins) by the microorganisms. It is in order next to point out that humus as a whole is characterized by certain specific physical, chemical, and biological properties that distinguish it from other forms of organic matter in nature.

Humus is practically insoluble in water, although some of it may go into colloidal solution in pure water. To a large extent it is soluble in dilute alkali, and certain of the humus constituents may dissolve in acid solutions.

One of the most important and characteristic properties of humus is its nitrogen content, which usually varies from 3 to 6 per cent, although the nitrogen concentration may be frequently lower or higher than these figures. The carbon content is usually 55 to 58 per cent. According to Waksman,<sup>1</sup> the average theoretical carbon content of soil humus is 56.24 per cent, and the average nitrogen content is 5.6 per cent. This gives a theoretical ratio of carbon to nitrogen of 56.24:5.6, or 10.04, which is very close to the one commonly found in the humus of soils. This ratio varies with the nature of the humus, the stage of its decomposition, the nature and depth of the soil, and climatic and other environmental conditions under which it is formed.

Humus is usually in a highly dynamic condition, for it is constantly being formed from plant and animal residues and is continuously being decomposed by microorganisms. It serves as a source of energy for various groups of microorganisms and with decomposition gives off carbon dioxide, ammonia, and other simple end products of decay.

Humus is highly colloidal and, as clay, it functions as an acidoid or micelle and usually carries a large number of negative charges. It is composed principally of C, H, O, N, S, and P, in contrast to the mineral acidoids, which are composed chiefly of Si, O, Al, and Fe.

<sup>1</sup> *Humus*, S. A. Waksman, Williams and Wilkins Co., 2nd edition, 1938, p. 182.

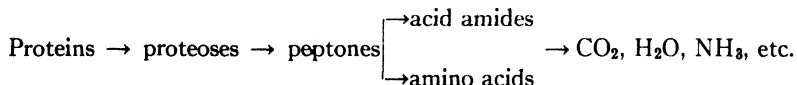
Humus is characterized by a high cation-exchange capacity; it combines with various inorganic soil constituents; it absorbs large quantities of water and exhibits the properties of swelling and shrinking. It does not exhibit so pronounced properties of adhesion and cohesion as do the mineral colloids and is less stable because it is subject to microbial decomposition. It has already been shown that soil humus is an important factor in the control of aeration, water-holding capacity, and granulation of field soils. Humus possesses other physical and physicochemical properties which make it a highly valuable soil constituent.

**Changes Brought about in the Nitrogen Compounds.** Most of the nitrogen added to soils by the plowing under of sod, crop residues, stable and green manures, and organic nitrogenous fertilizers is in the form of proteins and their derivatives. These forms of nitrogen are relatively insoluble. Since most of the nitrogen in the soil occurs as a constituent of the organic matter, the rate of its depletion from soils is decreased. For the most part, these organic nitrogenous substances cannot be used directly by higher plants but first have to be broken down into more simple compounds. The breaking down of any organic constituent added to soils is carried out by the action of microorganisms, such as bacteria, fungi, and actinomyces, and is in general in the direction of the formation of simpler forms from more complex compounds.

Plant and animal proteins and their derivatives in the soil break down into acid amides and amino acids<sup>2</sup> of various kinds, which in turn may be readily broken down to carbon dioxide, ammonium compounds, and certain other simple end products.

The final nitrogen product of the breakdown of nitrogenous organic material in soil is, chiefly, ammonia. The process of ammonia formation from nitrogenous substances is called *ammonification*. This ammonia may be used by plants or microorganisms, reduced by certain bacteria, leached out in drainage water, or oxidized to nitrates. The latter process is known as *nitrification*. In the utilization of these simple inorganic nitrogenous compounds by soil organisms, complex nitrogen compounds are again formed which in turn must be broken down by other organisms before the nitrogen becomes available to higher plants.

<sup>2</sup> Protein decomposition probably proceeds as follows:



**Changes in the Mineral Constituents of Organic Material as It Decomposes.** As the soil organic matter decays with the gradual formation of humus, simple mineral products begin to be released. Most of these simple end products are readily lost from the soil in the drainage waters, unless taken up by growing plants; and, if an adequate supply for normal plant growth is to be maintained in the soil, frequent additions of fresh organic matter must be made.

The principal mineral elements contained in organic matter are sulfur, phosphorus, potassium, magnesium, and calcium. Some are held in organic combinations, as are sulfur and phosphorus, and others are present in fresh organic materials as inorganic compounds.

As the decay of organic matter progresses, sulfur is left behind as a by-product of the general-decay types in the form of elemental sulfur or hydrogen sulfide or both. Some sulfur may be released as carbon disulfide. Elemental sulfur and hydrogen sulfide, upon oxidation, serve certain autotrophic bacteria as excellent sources of energy (p. 202). The biochemical oxidation of sulfur or hydrogen sulfide to sulfites and in turn to sulfates is called sulfofication. The oxidation of sulfites to sulfates occurs so rapidly that there is little chance for the accumulation of any great quantity of sulfite. It is thus observed that sulfur which was originally present in complex organic combinations is eventually released as a simple by-product and is made available to higher plants as the  $\text{SO}_4$  ion. This is the form of sulfur generally used by plants, and large amounts are lost in the drainage waters.

The soil organic matter carries considerable quantities of phosphorus which are rather readily liberated as the organic matter undergoes decay. The phosphorus is probably released as the  $\text{PO}_4$  ion, which in turn quickly reacts with other soil constituents and usually is readily fixed or held in the soil. As a result of this rapid fixation, which occurs in most soils, very little phosphorus is lost in the drainage waters. Since the quantity of inorganic phosphorus in soils is generally low and only slowly available, phosphorus added in organic matter is of great importance.

Other simple mineral products of organic matter decomposition that should be mentioned are potassium, calcium, and magnesium. They are released in ionic form and, under favorable conditions, react with water and in turn with  $\text{CO}_3^{2-}$  and  $\text{HCO}_3^-$  ions. Thus a certain portion of these mineral elements is again made available for a new generation of plants. This explains how it is possible for deep-rooted crops such as alfalfa and sweet clover to bring up considerable quantities of mineral

elements to the plow soil from the lower soil horizons, especially when these crops are used for green-manuring purposes.

In connection with the discussion of the changes in the mineral constituents accompanying organic matter decomposition, it should be recalled that considerable quantities of mineral elements, as they are released, may be assimilated by microorganisms. Thus there is always competition between microorganisms and higher plants for available plant nutrients, and the needs for the microorganisms must be satisfied first. However, under normal soil conditions the competition offered by the microorganisms for mineral elements is in general of no particular consequence. Where large amounts of carbonaceous materials are added to soils, a nitrate deficiency frequently occurs owing to the assimilation of the nitrogen by microorganisms active in the decaying of the organic matter.

**Organic Compounds as Direct Sources of Plant Food.** Over a hundred years ago the *humus theory* of plant nutrition was advanced by Thaer, a German. According to this theory, humus was absorbed directly and assimilated by higher plants. This theory was generally accepted until it was ridiculed by Liebig and replaced by his mineral theory. The mineral theory indicated that the mineral elements are the principal nourishment of plants, and Liebig believed that the main value of humus in soils was due to its release of ammonia into the atmosphere where the ammonia was absorbed directly by plants. The belief that organic matter is not used directly by higher plants was held from Liebig's time until rather recently. It is now generally believed that some organic substances do serve as suitable nutrient material and are used directly by higher plants without undergoing decomposition. These organic compounds are absorbed in a complex molecular condition, but apparently only a very small portion of the soil organic matter is absorbed as such.

A field of soil and plant research that recently has received considerable impetus is concerned with stimulants to plant growth. It is believed that they develop within the soil through microbial action or are produced by the plant itself. Many investigators believe these stimulants (frequently referred to as auximones, hormones, phytamine, or growth-promoting or growth-regulating substances) function directly in the nutrition of plants in a manner analogous to vitamins in animal nutrition. These substances affect rates of cell growth, root formation, and flower formation of certain plants, and they also favor the development of some groups of microorganisms. It must be added that comparatively little is known concerning the formation or liberation in the

soil of organic substances through the direct or indirect activities of microorganisms, which modify the growth of higher plants.

It must not be inferred that all organic substances released or produced in the soil are beneficial to higher plants. Under certain conditions several organic compounds are found in soils that are known to be harmful. It is generally believed that such substances are the result of improper soil conditions and are not the direct cause of decreased crop yields. When such unfavorable soil conditions are corrected, the organic toxins usually disappear rapidly as a result of microbial attack. Rational soil-management practices, such as providing good drainage and tillage and supplying lime and fertilizers when needed, usually prevent the accumulation of organic soil toxins.

### EFFECTS OF ORGANIC MATTER ON SOIL CHARACTERISTICS AND PRODUCTIVITY

Highly decomposed forms of organic matter, if present in appreciable quantities, have a marked effect on the chemical and physical properties of mineral soils. The beneficial effects of humus on the tilth of soils is more noticeable in those that are very sandy and those high in clay content than in soils of medium texture. However, a liberal quantity of humus aids materially in maintaining a desirable structural condition in all classes of soil. As was pointed out in Chapter 5, humus contributes greatly to the ion-exchange capacity of soils and this function is of special importance in soils low in content of clay minerals. The soil properties chiefly influenced by humus are suggested in the following questions.

#### Questions

1. How does organic matter affect physical properties of soil?
2. Do these physical changes influence productivity?
3. What chemical processes in soil are affected by humus?
4. Are beneficial biological processes influenced by organic matter?

**Physical Effects of Organic Matter in Soils.** Organic matter is of extreme importance in regard to crop growing because of the many useful effects it produces in soils, and many of these effects are purely physical. Organic matter increases the water-retaining power of soils, decreases water-runoff losses, improves aeration especially on the heavier soils, and produces a better soil structure or tilth. Owing to the fact that organic matter improves the tilth of soils by promoting granulation and to the fact that organic matter decreases water-runoff losses, the

damage done by either water or wind erosion is greatly reduced. Sandy soils often are subject to wind erosion, which can be controlled to some extent by increasing the soil organic content. Organic matter serves to bind sandy soils and lessen their movement by wind.

That portion of the soil organic matter which has undergone considerable decomposition assumes colloidal properties for the most part and as such has a very high absorptive capacity for water. On the dry-weight basis, humus has a water-holding capacity of several hundred per cent and may be considered to act much like a sponge. Soil water is also retained in the small pores or air spaces between soil particles. In the more sandy soils these spaces frequently are too large for maximum water retention, and humus tends partially to fill these large spaces and make them a more effective size for holding water. The humus also tends to pull the sand particles together, thereby increasing water retention. In heavy clay soils the pore spaces between the mineral particles frequently are too small for the greatest moisture storage. Organic matter improves this condition by forcing the soil particles apart, thus increasing the ability of those soils to retain water. The water-holding power of all mineral soils can be increased by raising the humus level.

**Plant Growth Increased by Good Tilth.** Soil tilth refers to the physical properties of a soil as related to crop growth. The term *tilth* is used frequently to denote the structure of a soil. A soil in good tilth is easily cultivated, is loose and mellow, and is characterized by being well granulated. The degree of granulation in all mineral soils is determined to a large extent by the quantity of soil organic matter. Sandy soils are not well granulated because they lack sufficient binding material to hold the particles together. Organic matter, added to these soils, tends to promote granulation by binding the sand particles into clusters, thereby improving the tilth of these soils. Heavy clay soils, low in organic matter, often are in poor tilth because of clodding. The thorough mixing of organic matter in these soils promotes granulation, and clodding may be greatly decreased. A heavy clay soil that contains ample organic matter is usually mellow, friable, and easily tilled. Organic matter makes heavy soils less sticky, enables them to receive, distribute, and hold water more effectively, and makes them easier for roots to penetrate.

The degree of aeration in heavy soils is determined to a large extent by the structure or tilth of the soil, and so indirectly aeration is improved by the presence of organic matter. Sandy soils frequently are excessively aerated, and the addition of organic matter to these soils tends to diminish excessive ventilation.

The dark color of soils is due largely to organic matter. It is known that dark-colored surfaces are better absorbers of radiant energy of the sun than light-colored surfaces; therefore, dark-colored soils would be expected to warm up faster than light-colored soils, which are similar in all other respects, including the moisture content. This would lead one to expect dark soils, all other things being equal, to have a somewhat higher average temperature, which would tend to promote a more rapid seed germination. It is obvious that under certain circumstances, as in early market gardening, a hastening of seed germination and a more rapid early growth of plants may be highly desirable. However, attention should be called to the fact that, if the higher organic content is associated with a higher moisture content, the effect of the dark color on soil temperature may be more than offset by the increased quantity of heat required to warm up the additional water.

**Chemical Functions of Organic Matter in Soils.** Organic matter improves the soil chemically by serving as a storehouse or supply of plant-nutrient elements. As the organic materials are decomposed, the plant-food elements contained therein are gradually released. Most of the soil's nitrogen supply exists in organic form. Nitrogen cannot be held in the soils in humid regions to any great extent except as it is held in organic combination. This organic nitrogen gradually undergoes conversion into nitrate under normal soil conditions, and in the absence of growing plants most of the nitrates may leach out. Decomposition of organic matter favors the release of plant-food elements from the soil minerals. Various organic and inorganic acids are produced in soils when organic matter decays (p. 223), and they have a very pronounced dissolving effect on soil minerals. One of the important end products of organic matter decay is carbon dioxide gas. This gas dissolves in the soil water and forms carbonic acid, which is an effective dissolving agent for the soil minerals; the dissolving effect of this carbonated water is several times that of pure water.

After the organic matter has undergone considerable decay, it usually assumes the colloidal state. The colloidal properties thus exhibited exert important physical and chemical effects which are directly concerned with soil productivity. The organic colloidal materials have a much greater cation-exchange capacity per unit weight than do the mineral colloidal materials, and hence they may act as buffers in the soil, thereby retarding the processes by which changes in soil reaction (acidity or alkalinity) are produced. Furthermore, these colloidal substances have a strong ability to adsorb or hold on to the constituents

of fertilizers and nutrients released from soil minerals, thus decreasing their rate of loss by leaching.

**Biological Effects of Organic Matter in Soils.** Organic matter improves the soil for the growth of microorganisms which, after all, are the agents whereby the plant-food elements of the soil are kept in circulation. It serves as a source of food and energy for the majority of the soil microorganisms.

The soil may be considered a factory operating to produce plant nutrients. Soil microorganisms may be considered the power or driving force in this factory, and the soil organic matter the fuel or energy for this power. The organic matter is burned to carbon dioxide, water, ash, and various other products, the nature of which is determined mainly by the degree of soil aeration. That the organic constituents constantly are being burned in the soil is shown by the continuous evolution or output of carbon dioxide.

The complex constituents of organic matter, upon being decomposed, are simplified, and nitrogen in the ammonia form is released and changed to the nitrate form. The energy stored in compounds of growing plants for the most part eventually is either used or released by soil micro-organisms whose activities within the soil make food elements available for a new generation of crop plants. Thus it can be said that, without the presence of organic matter to supply food and energy for the soil microbes, the plant-food elements of the soil could not be changed to usable forms.

Only the more important physical, chemical, and biological effects of organic matter in soils have been briefly discussed, but enough has been said to indicate definitely that organic matter exerts a controlling influence on the soil and its productivity.

#### ACCUMULATION AND LOSS OF SOIL ORGANIC MATTER

The stock of organic matter in our virgin (uncultivated) soils as taken over by the pioneers was an accumulation during many generations. Its collection in the surface layers of the earth began several thousand years ago. As the rocks and minerals of the earth's crust decomposed, mineral elements were made available to plants; and, as supplies of nitrogen in usable chemical combinations were produced from the store of nitrogen in the air, plants grew, died, and contributed their remains to the soil. Thus organic matter began to accumulate. As the supply of available plant nutrients in the soil increased, the accumulation of soil organic matter increased accordingly. This condition continued

until an equilibrium point was reached at which the accumulated organic matter held in combination most of the mineral nutrients that the soil could supply in readily soluble forms. Thereafter, the quantity of organic matter remained more or less constant because the supply of minerals became a limiting factor in the growth of plants. This so-called equilibrium level is determined to a large extent by (1) the climatic conditions (temperature and rainfall), (2) the nature of the vegetation, (3) the topography of the soil (which influences erosion losses), (4) texture of the soil, and (5) drainage conditions.

Almost all virgin soils, where there is sufficient rainfall, are kept covered with a mat of dead grass or leaves. Under grass vegetation the entire top growth is left annually on the soil surface to decompose while much of the root system decays within the soil. In forested areas a certain amount of leaf litter accumulates on the surface of the soil each year together with fallen twigs, branches, and some logs. A large part of the year's growth, however, does not return to the soil. A small part of the forest litter is carried into the soil by worms and insects, and some of the more highly decomposed portions are carried downward in suspension in water.

The conditions that favor the accumulation of organic matter in soils are, in general, the reverse of those favoring its decomposition. The accumulation of organic matter represents the difference between the addition through plant growth on the one hand and the decay activities of microorganisms on the other. Under natural soil conditions, an equilibrium exists between the formation of organic matter and its decomposition. The balance is determined largely by soil and climatic conditions. Cultivation disturbs this natural equilibrium because less organic matter is usually returned to the soil and decomposition processes are speeded up by farming operations. The accumulation and loss of organic matter may be considered by answering the following questions.

### Questions

1. Do climatic conditions influence the accumulation of organic matter?
2. How much organic matter do soils contain, and how is it distributed in the profile?
3. Is there a loss of organic matter when virgin soils are brought under cultivation?
4. What conditions favor organic matter accumulation?
5. What are the basic principles involved in organic matter accumulation?
6. How can organic content be maintained or restored?
7. Is it feasible to maintain a high organic content in soils?

**Climate a Factor in the Organic Content of Soils.** The quantity of organic matter in a soil represents the balance between addition and decomposition. Accordingly, climatic factors which affect the quantity

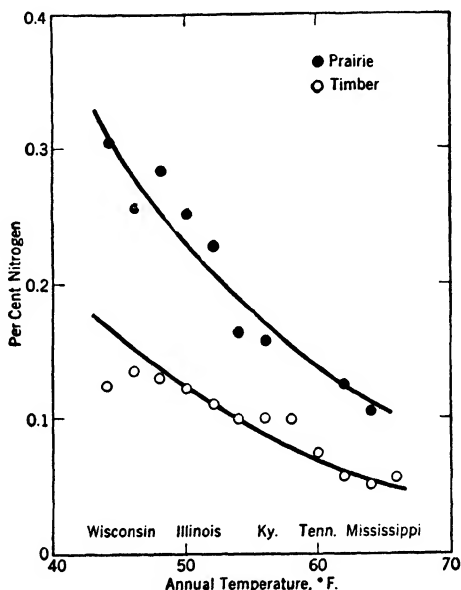


FIG. 48. Nitrogen-temperature relation in humid prairie (upper curve) and humid timber soils (lower curve) for silt loams. [Missouri Research Bull. 152.] These curves show that nitrogen and organic matter contents of soils under similar moisture conditions decrease from north to south, and that under similar conditions the prairie soils are considerably higher in nitrogen and organic matter than the timber soils. In these studies the average nitrogen or humus content of a county area was chosen as a comparable unit. All nitrogen analyses for a county were arranged according to topography, vegetation, and texture of the soils, and their respective averages were tabulated with the mean annual temperature and annual humidity factor of the county.

of organic material developed in or returned to the soil and the activity of decay organisms have a bearing on the amount accumulated. Studies have shown that, when average annual temperatures decrease and the moisture relations remain constant, the quantity of organic matter in soils of similar characteristics and covered by the same type of vegetation increases (Fig. 48). The increase is somewhat greater in grassland soils than in forest soils.

On the contrary a decrease in moisture supply with temperatures remaining constant results in a decrease in organic content in soils of similar characteristics and vegetative cover. Again the change is more pronounced in grassland than in timbered soils (Fig. 49).

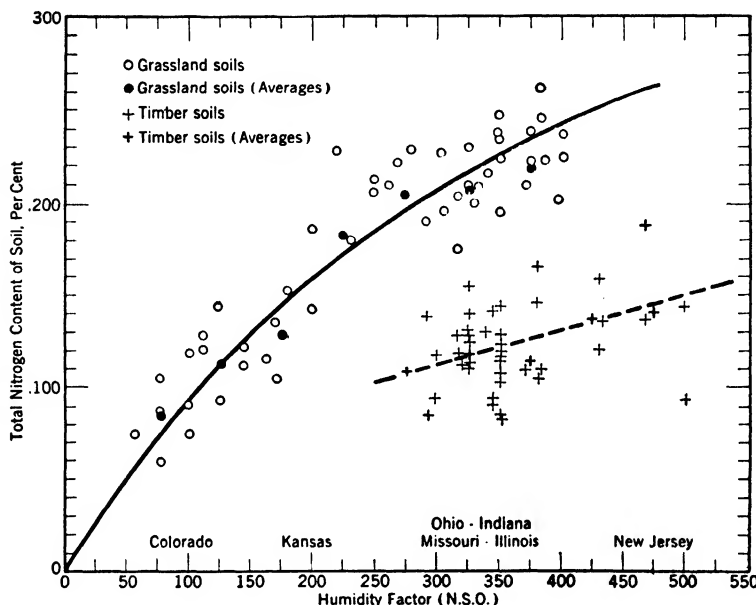


FIG. 49. Soil nitrogen-humidity factor relationship along the annual isotherm of 11° C. [Missouri Research Bull. 152.] It is observed that with an increase in humidity, with all other factors constant, including the mean annual temperature, there is an increase in content of soil nitrogen in both the grassland and timber soils. N.S.Q. is the ratio of precipitation to the absolute saturation deficit of the air. These values are used instead of precipitation values as such since rainfall alone is not a satisfactory index of soil-moisture conditions because of the great variations in evaporation. The N.S. quotients include therefore the effect not only of temperature but also of air humidity on evaporation.

In view of the long growing season and high rainfall of the Southeastern States, which results in a large amount of plant growth, one might expect the organic content of the soils to be correspondingly high. Such is not the case, however, because the long, warm, moist seasons are also favorable for decomposition, with the result that corresponding soils in the North are higher in organic matter than in the South. The decreasing organic content of soils as one travels westward into areas

of lower rainfall was pointed out in the study of the great soil groups.

**The Quantity and Distribution of Organic Matter in Representative Soils.** Although highly important in soils, organic matter makes up only a very small fraction of the total weight of mineral soils. The actual amount of humus found in any soil depends on texture of surface and subsoil horizons, topography, drainage, climatic factors, native vegetation, and treatment. The quantity ranges from a fraction of 1 per cent for the very poor sandy soils and desert soils to 12 or more per cent for prairie soils. The humus content is much higher in organic soils. Generally speaking, lighter soils (sands and sandy loams) are lower in organic matter than the heavier soils (clays, clay loams, etc.) because the sandy soils did not originally support as dense a vegetative growth, because they have better aeration which promotes more rapid decomposition, and because they are more subject to leaching losses.

In humid-region soils the nitrogen is carried or held almost entirely by the organic matter, and, as the content of organic matter varies, so does the nitrogen. The nitrogen content of soils in the United States, according to the results of analyses of a large number of samples carefully collected from selected areas under natural conditions, is given in Table 27. These samples are from six natural soil groups or regions. Each region contains soils that are extremely high and soils that are

TABLE 27  
AVERAGE NITROGEN CONTENT IN VARIOUS SOIL REGIONS OF THE  
UNITED STATES \*

| Soil Region    | Nitrogen, per cent |                                  | Pounds of N per<br>Acre to Depth of<br>40 Inches |
|----------------|--------------------|----------------------------------|--|
|                | Surface 6 inches   | Average to depth of<br>40 inches |  |
| Brown forest   | 0.05-0.20          | 0.05                             | 6,700  |
| Red and yellow | 0.05-0.15          | 0.03                             | 4,000  |
| Prairie        | 0.10-0.25          | 0.12                             | 16,000   |
| Chernozem      | 0.15-0.30          | 0.12                             | 16,000   |
| Chestnut       | 0.10-0.20          | 0.08                             | 10,700   |
| Brown          | 0.10-0.15          | 0.06                             | 8,000  |

\* "Soil Nitrogen," O. Schreiner and B. E. Brown, *U.S.D.A. Yearbook* 1938, p. 366.

extremely low in nitrogen, and obviously such figures can only approximate an average content, yet they are of interest and they present the best available picture of the quantity of nitrogen in the soils of the United States.

It is interesting to note that the nitrogen supply is highest in the soils of the Prairie and Chernozem regions and that the quantity declines both east and west of these groups. The approximate quantities of organic matter may be obtained for the different soil regions by multiplying the values in Table 27 by the factor 20.

The data presented illustrate, furthermore, that the humus is usually concentrated in the upper soil layer and diminishes rapidly in the subsoil, although it should be noted that in the Prairie and Chernozem soils the organic matter extends to a much greater depth than in the timbered soils. The general distribution of organic matter in representatives of several of the great soil groups is shown in Fig. 50. Since most of the organic matter is found in the topsoil, any erosion that occurs will result in significant organic matter losses. Furthermore, it is the topsoil that is disturbed by various tillage operations which greatly increase the loss in organic matter and nitrogen.

**Organic Matter Depleted by Cultivation.** Even on non-erosive land that is brought under cultivation, rapid losses of organic matter usually occur. It has been observed that the losses are most rapid immediately after farming is started, and thereafter the rate of disappearance is decreased; ultimately the organic content of the soil will reach a comparatively fixed level. Under all comparable conditions, this rather stable organic matter level is much higher for good systems of soil management than for poor ones.

It has been illustrated at the Missouri Agricultural Experiment Station<sup>3</sup> that, as a result of cultivation over a period of 60 years, soils in a non-eroded condition lost over one-third of their organic matter, the losses being much greater during the earlier than the later periods. The organic matter losses amounted to about 25 per cent the first 20 years, about 10 per cent the second 20 years, and only about 7 per cent the third 20 years. In other words, a somewhat fixed equilibrium was attained after about 30 years (Fig. 51). In dry-land farming, the losses may be even greater.

That the organic matter of the soil which receives little or no added material becomes depleted is evident from the fact that there is a con-

<sup>3</sup> "Soil Fertility Losses under Missouri Conditions," H. Jenny, *Missouri Agr. Exp. Sta. Bull.* 324, 1933.

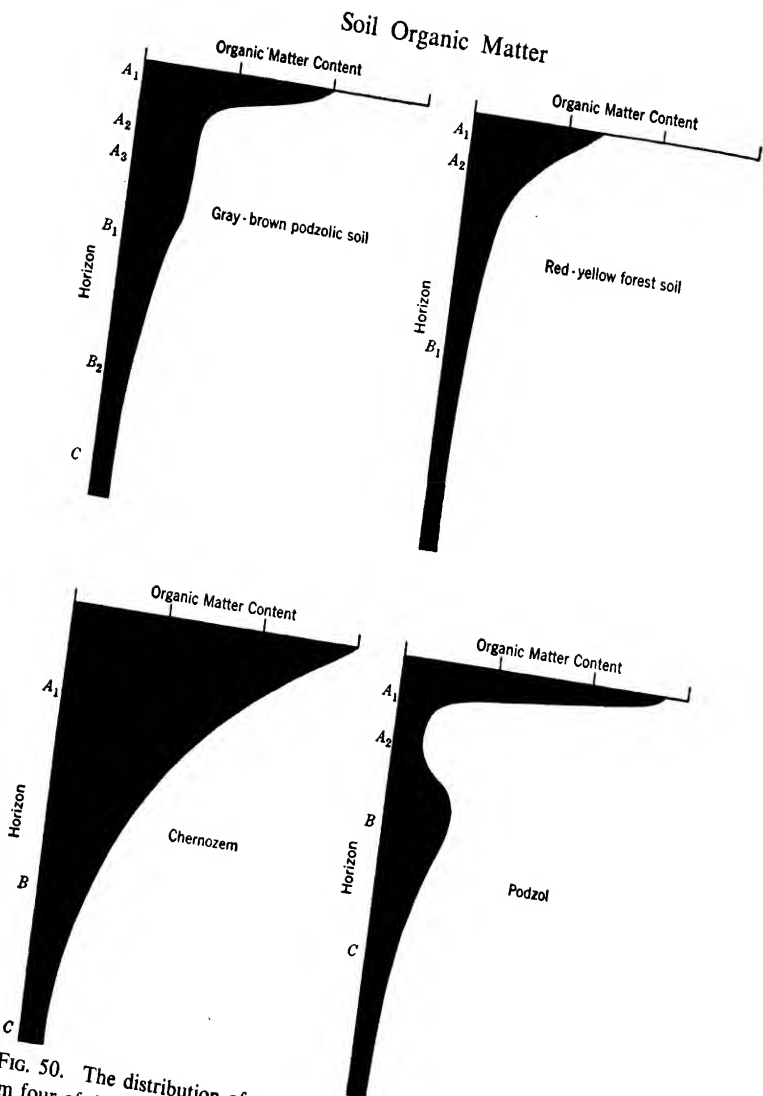


FIG. 50. The distribution of organic matter in the profiles of heavy soils from four of the great soil groups (generalized presentation).

tinuous evolution of carbon dioxide from soils under favorable moisture and temperature conditions. It has been estimated that a soil of average organic content under humid temperate conditions may lose 2.5 tons of organic matter per acre each year.

If a ton of fresh organic matter (dry weight), in the form of green manure, farm manure, or plant stubble, is plowed under or worked into

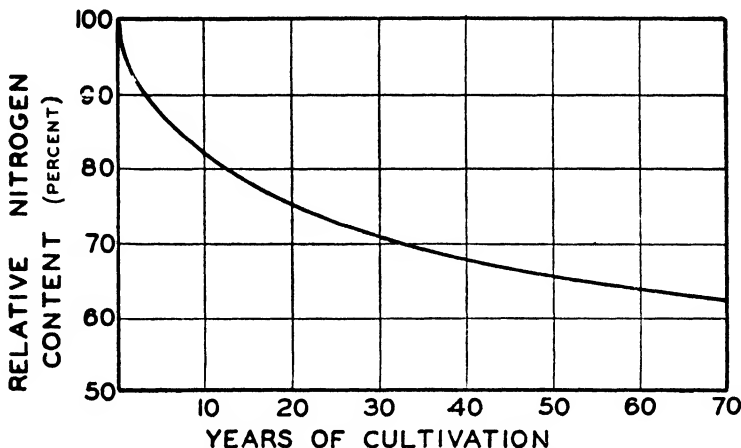


FIG. 51. Decline of soil fertility with length of cultivation period under average farming practices in the Middle West. [After Jenny, *Missouri Agr. Exp. Sta. Bull.* 324, 1933.]

the soil, decomposition begins immediately. Within a period of two or three weeks, under favorable conditions of moisture, temperature, and aeration, only about one-half of the original ton will be left. After a month or six weeks have passed, only about 500 pounds of the original ton may remain. Of course, the quantity remaining will vary greatly from one set of conditions to another, but the point of emphasis is that there is a continuous dissipation of the soil organic matter.

**Soil Conditions Favoring the Accumulation of Organic Matter.** The quantity of organic matter in soils and its rate of accumulation are determined by numerous factors, the most important of which are (1) the nature and amount of organic materials returned, (2) soil moisture, (3) soil temperature, (4) the degree of soil aeration (which is determined largely by the soil texture, tillage practices, and drainage conditions), (5) topography, and (6) the nature and supply of plant nutrients. All these factors except the first may be classed as soil factors. Of course, soil moisture and soil temperature are governed largely by the corresponding climatic factors.

Organic matter decay is essentially a burning process (carried out by microorganisms), and as in any burning process, the greater the supply of air, the more rapid will be the burning and the more completely will the organic matter disappear. This explains why cultivation tends to increase the rate of organic matter decomposition in the topsoil and emphasizes the importance of a minimum of cultivation on sandy soils.

Soil moisture and soil temperature have an important influence on organic matter accumulation. With ample moisture and high temperatures organic matter disappears very rapidly, especially from light-textured soils. With lower temperatures and heavier soils the disappearance is less rapid. Poor soil drainage aids in the accumulation of organic matter, but it must be remembered that the activity of the organic matter is perhaps more important than its mere presence.

In general, organic matter accumulation is greater in comparatively level soils than in soils of rougher topography, particularly if water erosion is in evidence. Since most of the organic matter is found in the upper soil horizons, any water erosion that may occur will result in significant organic matter losses. Frequently soils of level topography exhibit poor profile drainage or have a high water table, and either of these conditions may aid in the accumulation of organic matter.

The humus level in mineral soils is very closely associated with the supply of the nutrient elements such as calcium, nitrogen, phosphorus, and potassium. Many other elements, of course, are needed by plants, but these are the ones that commonly need to be considered in agricultural practices. Since plant growth is the source of nearly all the soil humus, the quantity of plant growth and hence the rate of humus formation depend to a large extent on the supply of available plant nutrients in the soil. For the maximum accumulation of organic matter, the soil must be well supplied with the essential plant-nutrient elements. Crops used for adding organic matter to the soil not only make more top growth but also make denser and more extensive root development. Many soils are too low in fertility to produce either a good sod, cover, or green-manure crop. With such soil conditions, it is utterly impossible to build up the humus content by attempting to grow soil-conserving or soil-improving crops. If the organic matter is to be restored in soils that have lost their organic matter and their soluble minerals as well, provision must be made for restoring these mineral deficiencies before attempting to grow crops for the sake of adding organic matter.

**Fundamental Principles Involved in the Accumulation of Soil Organic Matter.** The best practices for maintaining or increasing the organic supply in soils vary from one section of the country to an-

other or from one farm to another, and frequently vary from one field to another on any particular farm. It is logical, then, that this problem be discussed from the standpoint of the fundamental principles involved, and certain of them may be applied as they pertain to the particular conditions and circumstances concerned.

The accumulation of organic matter in soils is primarily a nitrogen problem. Theoretically, there can be no increase in effective soil organic matter without first a proportionate increase in soil nitrogen. This implies that there is a very constant and close relationship between the nitrogen and organic matter contents in soils. This close relationship does actually exist (p. 238). Since the ratio of C to N in humus is roughly 10 or 12:1, it must be concluded that neither carbon nor nitrogen, and hence soil organic matter, can be permanently or appreciably increased or decreased without a corresponding change in the other. Permanent soil productivity requires practices that maintain the soil nitrogen. It has also been pointed out that in humus the nitrogen is associated with lignin. It is essential, therefore, that lignin as well as nitrogen be present if humus accumulation is to result. Because the nitrogen content of most plant residues added to soil is low, and mature plant tissue is high in lignin, much emphasis is placed on nitrogen supply in the effort to increase the humus content of soil.

If the nitrogen content of plant residues is low, added nitrogen will be required to meet the demands of the soil organisms which produce the soil humus. We must therefore come to the conclusion that the accumulation or the restoration of soil organic matter is a problem of utilizing nitrogen as a means of holding carbon and other materials that constitute humus. Since nitrogen is an essential constituent of humus, the value of organic materials added to soils as a source of humus is in a general way directly proportional to their content of nitrogen. The carbon to nitrogen ratio of several organic materials is presented in Table 28.

**Maintenance and Restoration of Soil Organic Matter.** Although there is a rapid depletion of the soil organic matter immediately after virgin lands are brought under cultivation, there is some consolation in the fact that this high rate does not continue indefinitely. It has been emphasized that, after a period of heavy loss, a fairly constant level is attained during a long period of continued cultivation; the level is determined by the environment associated with a particular soil. Once the organic content has reached a low level, restoring the organic matter to its original level usually proves to be a rather expensive and an extremely slow process, because the producing power of these soils has

TABLE 28

## THE CARBON-NITROGEN RATIO OF SOME ORGANIC MATERIALS \*

| Material                 | C:N Ratio |
|--------------------------|-----------|
| Sweet clover (young)     | 12        |
| Barnyard manure (rotted) | 20        |
| Clover residues          | 23        |
| Green rye                | 36        |
| Cane trash               | 50        |
| Corn stover              | 60        |
| Straw                    | 80        |
| Timothy                  | 80        |
| Sawdust                  | 400       |

\* Data taken from several sources. The values are approximate only, and the ratio in any particular material may vary considerably from the values given.

been permitted to decline to a degree that makes it difficult to grow green-manuring or soil-improving crops without special treatment or fertilization.

In other words, many soils are too low in fertility to produce a good sod, cover crop, or green-manure crop. With such soil conditions it is utterly impossible to build up the humus content unless material from the outside is brought in, and that is usually impractical. If the organic matter is to be restored in the soils that have lost most of their organic content and their soluble minerals as well, provision must be made to replace these mineral deficiencies before attempting to grow crops for the sake of adding organic matter.

The humus level in mineral soils is very closely associated with the supply of the nutrient elements, such as calcium, phosphorus, potassium, and nitrogen. Since plant residues are the source of nearly all the soil humus, the quantity of plant growth and hence the rate of humus formation depend to a large extent on the supply of available plant food in the soil. For maximum speed of organic matter restoration, the soil must be well supplied with the essential plant-food elements in order that greater crop residues will be produced.

The rate of organic matter loss increases rapidly as the organic content is raised, and, as has been emphasized, its maintenance at a high level is not only difficult but also expensive. It is, therefore, usually unwise and uneconomical to maintain the organic matter above a level consistent with good crop yields. Attention, then, should be directed toward the frequent additions of small quantities of fresh organic materials rather than to practices of maintaining the organic matter content at any particularly high level.

Since it is the activity of the organic matter that gives it real value rather than the mere presence of it, the promotion of proper soil conditions in order that the organic matter may perform its normal functions is of great importance. Soil conditions should be the best. Drainage should be good in order to promote aeration and granulation. Lime should be added as needed, not only to give increased crop yields and permit the growing of legumes but also to increase the rate of organic matter decomposition by promoting biological activity and thus keep the organic matter cycle in more rapid progress.

In a consideration of the maintenance and activity of soil organic matter, three phases demand special attention. First, frequent additions of sufficient crop residues must be made; second, enough nitrogen must be supplied to permit a high organic matter level; and, third, an adequate supply of mineral nutrients must be provided for. Without doubt, one of the greatest limiting factors in crop production is the low humus supply or the low activity of the humus. The importance of nitrogen in this connection cannot be overemphasized. The organic matter content in soils can be maintained or restored and its activity promoted by good systems of soil management involving the proper use of cover crops, green-manure crops, manure, fertilizer, lime, and by providing adequate drainage. The problem of regulating the humus supply of soils is only one, though a very important one, of the many phases of the scientific management of soils.

**The Feasibility of Attempting to Maintain a High Soil Organic Content.** The maintenance of an adequate supply of organic matter in soil is rendered more difficult because of the large quantity that is dissipated each year. Since the rate of organic matter loss from soils increases rapidly as the organic content is raised, the maintenance of a high organic level is not only difficult but also expensive. Therefore, it is wise to maintain the organic matter at a level that will result in good crop yields. Obviously, this level will vary with the type of soil, with the rotation followed, and especially with climatic conditions. It is unwise to attempt to maintain as high an organic level in soils of the southern states as in the soils of the northern states. The higher temperatures in the South speed up decomposition processes and almost prohibit the maintenance of a high organic level. Under conditions which prevail over most of the eastern half of the United States, the most feasible practice is to make frequent additions of fresh organic materials with no special attempt to maintain a high organic matter level.

## Cover and Green-Manure Crops

Green-manure crops are those which are turned under for the purpose of improving the soil. Usually such crops are turned under when they are still green, thus the name green manure. Occasionally these crops may be allowed to mature or even to lie over winter before being plowed under and still may be classed as green manures. A cover crop is one used to cover and protect the soil, particularly over the winter. A cover crop aids in preventing erosion and leaching, protects the soil from the impact of rain, shades the soil, and aids in protecting the soil from excessive freezing and heaving. The term cover crop is usually applied only to crops planted for cover and soil-improvement purposes. Although the terms cover crop and green-manure crop are not identical, practically they are the same thing, and will be so considered in this discussion.

The use of green-manure crops is general in the Atlantic and Gulf states. Some of the most commonly grown cover and green-manuring crops in the southern states are Austrian winter peas, hairy vetch, lespedeza, crimson clover, and lupines. In the potato-growing areas of the humid regions of the United States, cover and green-manure cropping is frequently practiced, primarily to increase the organic matter supply of the soil. Rye, red clover, crimson clover, and alfalfa are the crops generally used. In the Corn Belt, sweet clover is the most popular green-manure crop before corn, the main object being to supply the corn crop with sufficient nitrogen throughout the growing season. Cover cropping in orchards is practiced in nearly all the fruit-growing sections of the United States.

It is generally conceded that there is ample opportunity for much improvement in soil conditions through the wider and more intensive use of cover and green-manure crops. In a discussion of this subject the following points are treated.

## Objectives

- A. The effects of cover and green-manure crops on soil properties.
- B. Principal cover and green-manure crops and their regional distribution.
- C. Utilization of cover and green-manure crops.
- D. Influence of turning under green manure on crop yields.

## THE EFFECTS OF COVER AND GREEN-MANURE CROPS

As a whole, the soils of the United States are not so fertile as they once were. Commercial fertilizers can be used effectively in adding plant nutrients to the soil, and farm manure is an excellent source of nutrients and organic matter. In few instances, however, is the supply of manure adequate to meet soil-improvement requirements. There is need to make a greater use of soil-management practices which will reduce loss of nutrients by leaching, control the devastating effects of erosion, and add organic matter to the soil. The use of green-manure crops will aid in these respects. Green-manuring is practiced in the expectation of increasing the yields of subsequent crops. In explaining how or why the turning under of crops has this effect, the following questions are answered.

## Questions

1. Can the organic matter and nitrogen content of soils be maintained or increased through the use of green-manure crops?
2. Do these crops aid in conserving soluble plant nutrients?
3. Does the use of green-manure crops tend to concentrate plant nutrients in the surface layer of soil?
4. Is the availability of plant nutrients in the soil affected by turning under green-manure crops?
5. How may green-manure crops improve the subsoil?
6. In what ways do green-manure crops protect the surface soil?
7. How are the activities of soil organisms influenced by turning under green crops?
8. What effect does green-manuring have on soil acidity?
9. What are the effects of green-manuring on the growth of fruit trees?

**Addition of Organic Matter and Nitrogen.** The amount of organic matter that may accumulate through the addition of green manures varies greatly with the amount and nature of the crop turned under, degree of soil aeration, and climatic conditions (Chapter 10). A large part of the plant material may decay and disappear as carbon dioxide

and water. In well-drained soils of hot climates this loss may be so rapid that no permanent increase in the soil organic matter is made even by turning under a heavy green-manure crop. Certainly no very large addition to the soil organic matter can be expected from turning under one or two green-manure crops. Because, owing to decomposition losses, only a part of the plant materials added becomes a constituent of the soil humus, it is obvious that a great many years would be required to raise appreciably the organic content of soils by green-manuring. By adding 2,000 pounds of dry matter per acre each year in the form of green manure and assuming that it is all permanently retained, 10 years would be required to raise the organic matter content of the surface soil by 1 per cent. This rather crude illustration brings out the fact that the main objective of green-manuring in most instances must be to maintain rather than to increase the quantity of organic matter in soils and to supply continually organic matter in a readily decayable form.

Since legume crops have the ability to utilize atmospheric nitrogen if they are properly inoculated, they will increase the nitrogen content of the soil when used as green manure. The amount of nitrogen so added depends on the kind of legume (Table 29), soil conditions, thick-

TABLE 29

THE AMOUNT OF NITROGEN TAKEN FROM THE AIR BY VARIOUS LEGUME CROPS \*  
(Pounds per acre per year)

| Investigator      | Place     | Crop                            | Total Nitrogen,<br>pounds |
|-------------------|-----------|---------------------------------|---------------------------|
| Arny and Thatcher | Minnesota | Alfalfa                         | 113 †                     |
| Arny and Thatcher | Minnesota | Sweet clover                    | 117                       |
| Whiting           | Illinois  | Alfalfa (4 crops)               | 132                       |
| Fred and Graul    | Wisconsin | Alfalfa                         | 64 †                      |
| Shutt             | Canada    | Clover, red                     | 75-150                    |
| Shutt             | Canada    | Clover, red (average, 10 years) | 51                        |
| Whiting           | Illinois  | Clover, red (4 tons)            | 106                       |
| Graul and Fred    | Wisconsin | Clover, red                     | 100                       |
| Duggar            | Alabama   | Vetch, hairy                    | 79                        |
| Whiting           | Illinois  | Cowpea                          | 86                        |
| Breal             | France    | Bean                            | 87                        |
| Fred              | Wisconsin | Soybean (25 bushels)            | 57                        |
| Fred and Graul    | Wisconsin | Soybean                         | 108                       |
| Albrecht          | Illinois  | Soybean and cowpea              | 107                       |

\* Taken from Table 21, *Root Nodule Bacteria and Leguminous Plants*, Fred, Baldwin, and McCoy, University of Wisconsin Press, p. 218, 1932.

† Average figure.

In several cases the high figures represent the gain in nitrogen from the growth of two or three crops.

ness and height of the stand, and stage of growth at which the legume is turned under (Figs. 52 and 53). The quantity of nitrogen in a legume crop includes what it has taken from both the soil and the air, but only that nitrogen taken from the air represents a true addition to the soil when the crop is turned under. It is generally assumed that on the average about two-thirds of the nitrogen in a legume is taken from the air and one-third from the soil.

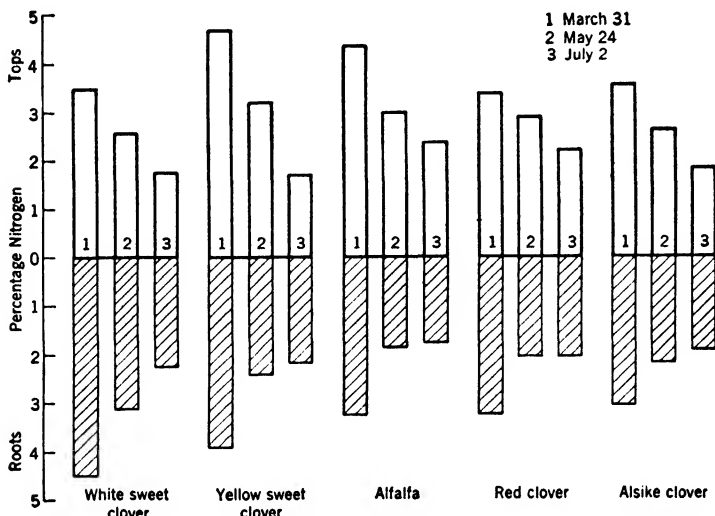


FIG. 52. Percentage of nitrogen in roots and tops of legumes at different stages of maturity. [*Ohio Agr. Exp. Sta. Bull. 405.*]

A generalized comparison of the distribution of dry matter and nitrogen in the tops and roots of several leguminous crops is shown in Figs. 53 and 54. It is observed that the annual legumes have comparatively small root systems in proportion to the mass of tops and that the percentage of the total nitrogen of the plant which is contained in the roots is less. The proportion of the plant nitrogen which is in the roots of annual legumes ranges from 5 per cent to 20 per cent, and that in the roots of the biennial or perennial plants from 24 per cent to 35 per cent. An acre of well-inoculated legumes, growing under favorable conditions, will perhaps gather 50 to 100 pounds of nitrogen from the air per year. If the amount of nitrogen in the soil is increased 50 pounds per acre, it is equivalent to over 300 pounds of sodium nitrate. The nitrogen so added is important not only as a nutrient but also as a means

of maintaining or increasing the humus level. The relationship of nitrogen and carbon ratios to humus formation are discussed elsewhere.

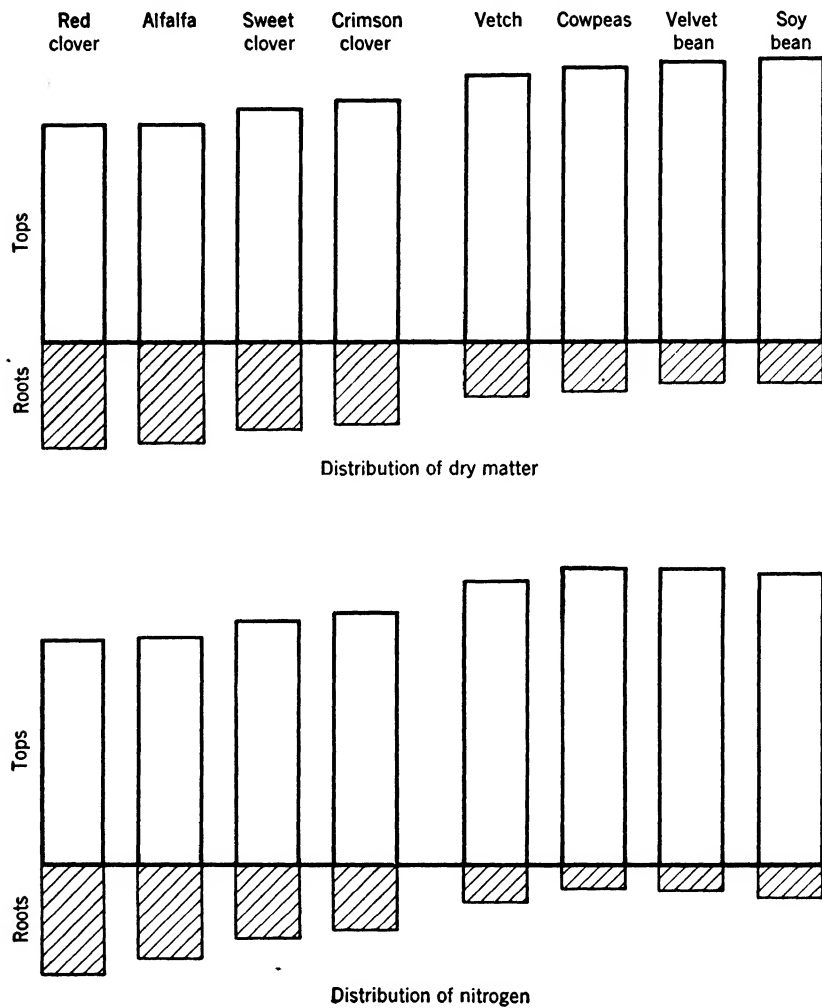


FIG. 53. The distribution of dry matter and nitrogen contents of several legumes between the tops and roots. [Calculated from data in *Root Nodule Bacteria and Leguminous Plants*, Fred, Baldwin, and McCoy, University of Wisconsin Press, 1932, p. 221.]

**Conserving Plant Nutrients.** Leaching losses of plant nutrients from fallow soils are much greater than from similar soils occupied with crops. Land lying idle after the removal of a crop during summer

or early fall is also often subjected to severe leaching losses. The nitrates are of particular concern in this connection, for they are highly soluble and are not adsorbed to any extent by the soil colloids. In the presence of a cover crop the soluble plant nutrients are taken into the plants and so are in little danger of being lost. Crops having extensive root systems are especially efficient in collecting plant nutrients, and thus they decrease leaching losses. When these crops are turned into the soil, decomposition takes place and the nutrients are released for a new generation of plants.

It is to be emphasized that the mineral content of a soil will not be increased by green-manuring although it may be conserved by decreasing wastage. The only elements actually added to the soil supply by green-manuring are carbon, hydrogen, oxygen, and (in legume crops) nitrogen.

**Concentration of Plant Nutrients in Soil Surface.** Green-manure crops, especially those which are deep-rooted, such as alfalfa, gather considerable quantities of plant nutrients from the subsoil. When the crop is turned under and decomposes in the upper layers of the soil, the nutrients as liberated are concentrated within a more limited area than previously. This permits the use of nutrients by succeeding shallower-rooted crops which otherwise would not have been available.

**Increased Availability of Plant Nutrients.** The addition of fresh organic matter to soils results in increased availability of certain inorganic constituents of the soil particles. The solubility of calcium, phosphorus, potassium, magnesium, and other elements is increased through the effect of the organic and inorganic acids produced as a result of the

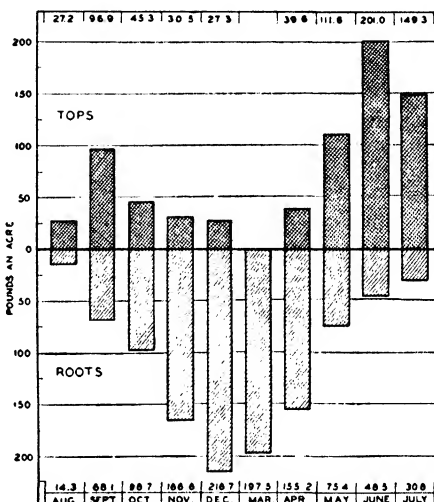


FIG. 54. Total nitrogen distribution in tops and roots of sweet clover at different dates during the year. "Although the distribution of nitrogen between roots and tops varies greatly with the season, the total nitrogen of the sweet-clover plant increases little after growth ceases in the fall of the first year." [From *Illinois Exp. Sta. Bull.* 394.]

decomposing organic matter. These increases are obtained in addition to the mineral constituents contained in the organic matter itself.

A greater availability of certain nutrients may be brought about in another way. The action of growing roots of certain plants, used for green-manuring purposes, may be more effective in utilizing a particular element in the soil than the roots of a following crop grown for harvest. This plant-food element is taken into the plant and returned to the soil in the green manure in forms more available than formerly to succeeding crops. Sweet clover, for example, may be able to take in phosphorus which is not available to some other crops.

**Improvement of the Subsoil.** Plants having long tap roots penetrate the subsoil deeply where possible, and, when the roots decay, numerous channels are formed which facilitate the circulation of air and the percolation of water. Furthermore, the nature of the subsoil will be modified somewhat through the increased quantities of organic matter resulting from the green-manure-crop roots, although no pronounced effect can be expected because of the limited extent of the root system in the subsoil.

**Protection of the Surface Soil.** Green-manure crops may serve as cover crops in that they may be used in many soils to protect the surface. On sloping land the soil is protected from water erosion, and light soils (both level and rolling) are protected from wind erosion. Furthermore, cover crops decrease the impact of beating rains which destroy the structure of the soil surface. In occupying the land over winter they hold snow, keep the soil warmer, and decrease root injury to perennial plants. Root injury may result from either alternate freezing and thawing or excessive cold. This may be very important in certain fruit-growing areas. Green-manure crops shade the soil to some extent and thereby decrease the loss of moisture by evaporation.

**Effect on Soil Microorganisms.** Organic matter supplied by green manures serves as food for many soil organisms and tends to stimulate certain biological changes to a marked degree. Such biochemical processes as the production of carbon dioxide, ammonia, nitrites, nitrates, other end products of decay, and the non-symbiotic fixation of nitrogen are all influenced by the incorporation of green-manure crops in the soil. As indicated above (p. 265), the increased availability of the inorganic constituents of the soil is a point that should be considered in this connection.

The effects produced by green manure on the microbial population depend not only on the kind of green-manure crop and its stage of

maturity but also on soil conditions such as drainage, lime content, and soluble nutrients. Young, succulent green manures are relatively high in nitrogen and, when turned into the soil, result in an immediate and marked increase in growth and activity of the microorganisms. Such green manures decay rapidly, and usually there is little delay in the accumulation of nitrates and the release of other simple end products of organic matter decomposition. On the other hand, when more mature crops having a low nitrogen content are turned into the soil, the organisms multiply rapidly and compete with higher plants for the available nitrogen. The nitrates may all disappear from the soil, and none will appear until after the rapid period of decomposition is past. Other things being equal, the lower the nitrogen content of the green-manure crop, the greater will be the delay in the accumulation of nitrates.

**Effect on Soil Acidity.** When a green-manure crop low in nitrogen is added to the soil, the first stage of decomposition may result in the formation of various organic acids, especially in the absence of free calcium carbonate. If soil conditions are favorable for further decomposition, the organic acids are broken down and the acidity disappears with the formation of carbon dioxide, water, and carbonates. On the whole, experimental results show that green manures generally do not increase soil acidity; if they do, such a condition is only temporary. With good soil-management practices green manures will have no lasting effect on increasing the lime requirement of soils.

**Effect on Growth of Fruit Trees.** The influence of cover and green-manure crops on the growth of fruit trees depends, among other factors, on the nature of the cover or green-manure crop, time of planting, locality, kind of trees, and soil conditions. These crops compete with trees not only for plant nutrients but also for water. In soils low in nitrates the competition for nitrates may become serious. Deep-rooted crops such as alfalfa and sweet clover draw heavily on the soil-moisture supply; and, at a time when the demands of the trees are great and the moisture supply is limited, this effect is of much concern. However, in soils that contain nitrate nitrogen in such quantities as to produce a prolonged, succulent growth of new wood, susceptible to winter injury, cover and green-manure crops serve to use the surplus nitrogen, slow up the growth of trees late in summer, and induce earlier maturity in the wood. Furthermore, crops that cover the ground during the winter hold snow, protect the tree roots from freezing, and on some soils help dry the soil in early spring.

## THE PRINCIPAL COVER AND GREEN-MANURE CROPS AND THEIR REGIONAL DISTRIBUTION

Green-manure crops may be classified or grouped in various ways. On the basis of their time of seeding and their occupation of the land, four types are recognized: (1) those occupying the ground exclusively throughout the summer; (2) those seeded with a nurse crop; (3) those seeded on grain stubble after harvest; and (4) those serving as winter cover crops. Certain legume and non-legume crops fit into and may be used in each of the four categories given. For convenience, these crops will be discussed under the two groups—legumes and non-legumes—and the following questions will be answered.

### Questions

1. What legume crops are commonly used for green-manuring purposes, and in what regions of the United States are they particularly adaptable?
2. Which of the non-legume crops are frequently used for cover and green-manuring purposes?

**Legume Green-Manure Crops.** Plants belonging to this group are commonly known as nitrogen gatherers. These plants, if properly inoculated and if grown under favorable soil conditions, contain nitrogen obtained from the air. When these crops are incorporated in the soil, they add nitrogen to the soil's previous supply. Several legume crops are grown in the United States for partial or complete use as green manures; among them the following are probably of most practical interest: hairy vetch, Austrian winter peas, crimson clover, crotalaria, sweet clover, alfalfa, cowpeas, soybeans, red clover, alsike clover, white clover, lespedeza, peanuts, Canada field peas, kudzu, lupines, bur clover, mung beans, and velvet beans.

Two distinct forms or varieties of hairy vetch are commonly sold. One is distinctly hairy, whereas the other is almost smooth. The hairy form is more winter-hardy but will not grow at as low a temperature as will the smooth form. Therefore, the former is preferred for planting in the more northern latitudes because of its winter-hardiness, and the smooth variety is preferred for planting in the South as a winter cover crop.

Austrian winter pea is the most winter-hardy of the field peas. It grows at quite a low temperature and in this respect is superior to hairy vetch. It makes a good growth under the winter conditions of the South

and is one of the best legumes for general use. There are several other species of winter peas used in some parts of the South.

Crimson clover makes an excellent winter cover crop for the northern part of the Cotton Belt, although stands are somewhat difficult to obtain. This has greatly limited its use. The seeds germinate easily, but the sprouts are readily injured by drying. A light rain may germinate the seed, but the young shoots may be subsequently killed by dry weather that often prevails at seeding time in the fall. Crimson clover is frequently seeded with a small grain or rye grass for winter grazing.

Lespedeza is finding much favor as a forage and soil-improving crop in certain sections of the South. Probably the Korean and Kobe lespedeza are the most generally used. A mixture of lespedeza and winter grain is frequently grown, and the lespedeza is used for green manure or hay after seed harvest. Georgia is growing about 1,000,000 acres of such a mixture.

Kudzu is filling a great need in southern states. It is used mainly on steep slopes and areas of what might otherwise be classed as waste land. Because of its vine-like habit of growth it is a good soil-stabilizing crop, and when well established it furnishes much hay and pasture. Poorly drained or waterlogged soils and soils with a hardpan closer than 20 inches to the surface are well adapted to the crop. Some idea of the rapidity with which the growth of this crop has increased may be gained from the fact that in Mississippi the acreage increased from 1,185 to 7,025 between the years 1939 and 1943.

Crotalaria is one of the most recent and important soil-improving crops in the South. It is well adapted to sandy soil and is better suited to a soil of low fertility than many other crops. The fact that the seed can be produced locally and at a low cost makes it an especially useful crop in the Cotton Belt.

Alfalfa is seldom used exclusively for a cover or a green-manure crop; yet when used in rotations it is recognized as one of the most important soil-improving crops. It is a deep-rooted perennial and improves drainage and aeration in heavy soils. It furnishes large quantities of organic matter. It makes an effective permanent cover and is recommended for orchards, where moisture is not a limiting factor, or for other permanent plantings in regions where it succeeds. It is grown as a cover and a green-manure crop in apple orchards in parts of Washington and Oregon and in certain irrigated sections of the West. In many parts of the North Central States, alfalfa is commonly used in crop rotation. The crop is either pastured or used for hay for two or three years, sometimes longer, and the aftergrowth is turned under for green-manuring purposes.

There are both annual and biennial forms of sweet clover, although the annual type is grown to only a limited extent in certain portions of the Corn Belt. The biennial type is used only in rotations and is perhaps the most satisfactory legume in the Corn Belt for use as a cover crop and as a green manure. It fits in well with a corn-wheat rotation. Sweet clover, like alfalfa, has a deep root system and produces a large tonnage of organic matter. These crops can be grown successfully only where the soil is well supplied with lime and phosphorus.

The various legume green-manure crops mentioned can be grown to best advantage only in definitely limited areas. However, the best crop to use in a given district must be determined finally by local conditions and by the objectives for which the green manure is grown.

In the northern parts of the United States, red clover, mammoth clover, alsike clover, field peas, common vetch, sweet clover, and alfalfa are commonly used. In regions having a moderate to warm temperate climate, cowpeas, crimson clover, red clover, sweet clover, and soybeans serve as green manures; and in the South crotalaria, Austrian winter peas, lespedeza, vetch, soybeans, cowpeas, crimson clover, red clover, kudzu, peanuts, mungbeans, and velvet beans are commonly recommended for use as green manure.

**Non-Legume Green-Manure Crops.** The chief difference between legume and non-legume green-manure crops is that legumes add both organic matter and nitrogen to the soil, whereas non-legumes add organic matter only. From the standpoint of maintaining the organic matter content of the soil, quantity or bulk is of great importance, and this can often be supplied more efficiently by such a crop as rye, sorghum, or oats than by a legume. It is to be remembered, however, that although a non-legume will add organic matter to the soil, the soil microorganisms may require additional nitrogen to effect rapid decay. The turning under of mature non-legume crops frequently results in a decrease in the yields of the subsequent crops because of the exhaustion of available soil nitrogen. If the nitrogen contained in the green-manure crop is not sufficient, the decay organisms will draw on the available soil nitrogen.

The principal non-legume green-manure crops are rye, oats, millet, buckwheat, rape, rye grass, sudan grass, corn, sorghum, and weeds. Rye and oats are perhaps more extensively used than any of the other crops listed. They make a rapid, abundant, and succulent growth, and they easily fit into almost any rotation. These crops are hardy and will grow under rather adverse weather and soil conditions. They are particularly useful on poor soils, but their beneficial influence is greatly

increased by seeding a legume crop with them. Combinations of rye and vetch and of oats and peas are very satisfactory for green-manuring purposes.

For the most part these non-legume crops which are used frequently as green manures are adapted to a wide range of soil and climatic conditions. They are not so exacting in their requirements as are many of the legumes. Most of these non-legume crops may be grown successfully throughout the humid regions of the United States. Rye is extensively used as a green-manure or a cover crop along the Atlantic seaboard; rye and winter oats are popular in the South; rye or rye and hairy vetch are successfully used in the North; and buckwheat is grown more often on the poorer soils of the North for green-manuring purposes than in any other section.

Weeds often serve as green-manure plants and are sometimes encouraged to grow for that purpose. Many orchardists allow weeds to grow in late summer, and they often make an effective cover crop. In certain tobacco sections weeds sometimes are encouraged in order that they may be turned under for green manure. In most instances it is believed that weeds are better than no cover or green-manure crop at all.

### THE UTILIZATION OF COVER AND GREEN-MANURE CROPS

The extensive use of cover and green-manure crops in the United States is a rather recent development and has coincided with the realization that land under continuous cultivation often rapidly loses its productive capacity. Green-manuring is only one of several practices used in the attempt to maintain soil productivity and is usually one of the last methods to be resorted to. To be most effective and to avoid detrimental results to soils and crops, the use of green manures should be well planned and the practice judiciously managed. In the practical utilization of cover and green-manure crops a number of problems, as indicated by the following questions, present themselves.

#### Questions

1. Under what conditions should green-manuring be practiced?
2. What points should be considered in selecting green-manure crops?
3. What precautions should be observed in preparing the seedbed, in seeding and fertilizing green-manure crops?
4. When should green crops be plowed under?
5. What crops are usually best to follow green-manure crops?

**Conditions under which Green-Manuring Should Be Practiced.**

When green manures are used properly they are highly effective, but when used injudiciously they may be useless or even harmful. They can generally be used profitably in soils that are low in nitrogen and organic matter because such soils usually possess unfavorable physical properties. Green manures are particularly beneficial on many light sandy soils and on heavy clay soils. They are quite generally used to advantage in orchards. On very fertile soils in good physical condition it is necessary to use green manures only at intervals sufficient to maintain the organic matter content, and this can usually be done by using a legume hay crop as part of the regular rotation. On livestock farms it is often most profitable to grow those crops which, like alfalfa and clover, can be utilized largely for forage, and to depend on the roots and stubble of these crops together with manure for the maintenance of the organic matter and nitrogen supply.

In general, green-manuring in some form should be practiced on most farms of humid regions and on irrigated lands elsewhere. This does not suggest a curtailment of the production of regular crops, but it does mean that cropping systems should be arranged so that a green manure may be worked in between regular crops. In other words, it usually is not advisable to sacrifice a summer crop for the purpose of growing a green crop to plow under because the value of the increased yields of following crops seldom equals the value of the crop plowed under. Summer green-manuring, however, usually can be recommended on very poor soils where drastic measures are necessary to begin the soil-improvement program. The same principle may apply to extremely light or heavy soils to improve their physical properties. A summer green-manure crop may also be used in orchards without interfering with the main crops.

The use of green manures in dry-farming areas in arid and semi-arid regions is, as a rule, impracticable. Not only is it difficult to produce such crops, but also they decay very slowly because of insufficient moisture. Not only do the green-manure crops require large quantities of water for their production, but also, when they are turned into the soil, evaporation losses are enhanced, owing to the increase in the amount of air space within the soil.

**Selection of Green-Manure Crops.** In choosing crops which are to be used for green-manuring purposes it is obvious that those crops should be used which are best adapted to the climate, soil, type of farming, and special purpose for which they are selected. Legume crops should be given preference when they can be grown satisfactorily, be-

cause of their ability, when properly inoculated, to add nitrogen to the soil. Difficulty in obtaining a catch of certain legumes and the relatively high cost of the seed are frequently determining factors in the choice of a legume.

Some of the legumes are annuals, some are biennials, and others are perennials. Some of them are warm-weather crops, and others grow well in cool weather; some are best adapted to fall planting and others to spring planting. The biennials and perennials as a rule make a slow growth at the beginning, and this is a serious disadvantage where quick results are needed. Alfalfa is a perennial, whereas sweet clover (the variety commonly grown) is a biennial. Red clover, alsike, and mammoth clover, as usually handled, may also be considered biennials. Cowpeas, soybeans, common vetch, field peas, peanuts, velvet beans, lespedeza, and crotalaria are summer annuals; hairy vetch is a winter annual in the North, and crimson clover, common vetch, and field peas are winter annuals in the South.

Where there is a choice between two or more crops, there is the question of which is preferred. In arriving at a decision, the following points should be considered: (1) the value of the crop as forage; (2) its ability to add nitrogen; (3) cost of seed; (4) ease of plowing under and of killing the crop; (5) the rooting habit of the plants; (6) rate of growth; (7) abundance and succulence of growth; and (8) soil requirements.

As a rule it is advisable to choose a green-manure crop that grows rapidly, one that produces abundant and succulent tops, and one that has the ability to grow well on poor soils. Cover and green-manure crops that interfere least with the regular cropping system should be used. In the North, catch crops following grain crops or planted with them are often used to advantage. On suitable land, sweet clover may be seeded with grain in the spring and turned under for corn the following spring. This is a common practice in the Corn Belt area. In the South winter cover crops fit into most farm programs effectively. In truck farming, catch crops can be used, and legume crops in the regular rotation may serve as green manure. In general farming no green-manure crop which requires an entire season for growing is likely to become widely used or generally profitable. Therefore, as a rule, it is necessary that crops be chosen which can be produced without the loss of the use of the land for an entire season. Such crops are often called "fill-in" or "catch" crops. For example, a crop such as rye may be seeded in the fall after corn has been cut for ensilage and plowed under for a second corn crop the next spring. In suitable climatic areas, soybeans may be seeded after oats or wheat and the crop turned under that

fall. The second crop of red clover may be plowed under when the setting of seed is poor and when the clover is not needed for pasture. However, the grower is perhaps justified in giving over the entire crop season to a green-manure crop when the succeeding crop is more or less permanent, as in orchard plantings.

When green-manure or cover crops are used in orchards, the season of growth of the trees must be considered. If an annual crop is used, it is necessary to plant it at a time that will offer as little interference as possible with tree growth, fruit development, and harvesting operations.

**Preparation for Green-Manure Crops.** Most of the green-manure crops are easy to grow under proper climatic conditions, even on relatively poor soils; nevertheless, if they are to be grown most successfully certain precautions should be observed. Consideration should be given to the preparation of the seedbed, the need for fertilizers, the need for lime, and the matter of inoculation for legumes.

As a general rule, more attention should be given to the seedbed preparation for legumes than for non-legumes. It is best that the soil be in good tilth and that a firm seedbed be established. Obviously, little or no seedbed preparation is necessary when cover or green-manure crops are seeded immediately after the removal of some cultivated crop, and, if seeding is made in a nurse crop that has already been planted, little can be done to modify the existing seedbed.

Many soils are too low in fertility to produce a good cover or green-manure crop. With such soil conditions, it is utterly impossible to build the humus content by attempting to grow soil-conserving or soil-improving crops. If the organic matter is to be restored in these soils by green-manuring, provision must be made to supply these mineral deficiencies before attempting to grow green-manure crops. In most soils where fertility has declined to a rather low level, a complete fertilizer—one carrying nitrogen, phosphorus, and potassium—is needed for non-legume crops. Legumes, if inoculated, usually will not require nitrogen fertilization. If fertilizer is needed in order to grow a good crop of green manure, it should be used; not only will the crops make more top growth, but also they will have a more dense and extensive root development. The plant nutrients in the fertilizer that are taken up by the green manure are not wasted because they will be available to the following crops when the green-manure crop decays. The kind and amount of fertilizer to use will depend primarily on the crop and soil conditions.

An abundant supply of lime is necessary for the maximum growth of legumes. The majority of soils contain sufficient lime for most non-

legume green-manure crops, but the supply frequently is inadequate for legume crops. Liming not only results in greater top and root growth but also increases the amount of nitrogen fixed by the legume bacteria. The relationship of the lime supply in soils to humus formation is discussed in Chapter 10.

It is emphasized again that, if legume green-manure crops are to be effective in increasing the nitrogen supply of soil, they must be well inoculated with an active strain of the proper bacteria. Most of the legumes require a particular kind of bacteria. For example, bacteria which produce nodules on soybeans will not do so on the alfalfa plant (p. 220). Inoculation may be accomplished by using artificial cultures. When one doubts if inoculation is needed, it is advisable to inoculate because the cost is relatively insignificant.

**Plowing Under Green-Manure Crops.** The time of turning under green-manure crops is an important consideration and is dependent on, among other conditions, the seasonal growth of the crop, the nature of the soil, the maturity of the crop, the weather conditions, and the crop which is to follow. Precautions in turning under green manures are of more concern if a crop is to follow immediately. Generally speaking, they should be plowed under when they are still green and succulent so that they will decay rapidly. This is particularly important in light soils.

A green-manure crop may remove much water from the soil, and, when the crop is turned under, the soil may be left too dry for the crop which is to follow. The decay processes which break down this organic matter are hindered, and this material will lie in the soil for some time without decomposing. When a large amount of organic matter is turned under, it may leave the soil so loose that the entire furrow slice will dry out and thus injure the following crop. Thus, when a crop is to follow immediately, it is better to plow the green-manure crop under some time in advance—while it is green and succulent and before the soil has become dry. If possible, green manure should be incorporated in the soil at a season when there is a plentiful supply of rain, for this will usually insure a rapid and effective decomposition. When large quantities of green organic matter are incorporated with the soil, decomposition products may cause considerable injury to the seedlings of the following crop. In order to avoid this type of injury, some time, perhaps two or three weeks, should elapse before planting.

A deficiency of soluble nitrates is likely to occur where green-manure crops are permitted to reach an advanced state of maturity before they are turned under, especially non-legume crops. The carbon-nitrogen

ratio is wide, and, in breaking down this carbonaceous material, the decay organisms may utilize most of the soluble nitrates in the soil in their life processes. The result is that the next crop may lack not only moisture but also available nitrogen. The younger the green-manure crop and the more water it contains, the shorter is the period of nitrate depression.

The effect which the time of turning under a green crop may have on leaching losses should also be considered. There may be appreciable leaching of nitrates when a green crop is turned under in the fall for a spring-seeded crop. This point should be given due consideration when a choice is to be made about fall versus spring plowing. As a rule, in the northern states plowing under a green crop in the spring generally gives best results. Where erosion or leaching is a problem, it is better to leave a green crop on the soil over winter and to plow in the spring. In the South, from the standpoint of the leaching loss of nitrates, green-manure crops should be turned under in the spring.

In plowing under green crops, it is best to distribute the organic material throughout the plow soil instead of having it in a more or less continuous layer at the bottom of the furrows. This may be accomplished, to some extent, by turning the furrow slice partly over instead of turning it over flat. When the amount of material to turn under is large, it is sometimes advisable to double-disk the land before plowing in order to cut up the material and to facilitate working it into the soil.

Obviously, no definite dates can be given for the turning under of a green-manure crop. There are certain precautions to be taken into consideration in relation to turning under these crops, but on the whole little difficulty is encountered and the farmer merely needs to use good judgment to secure good results.

**Crops to Follow Green-Manure Crops.** In general, it is a good plan to follow green manures with cultivated crops such as corn, tobacco, cotton, and potatoes. The cultural practices used in growing such crops produce conditions that favor decomposition of the green material and the consequent increase in available plant nutrients. Of the small-grain crops, rye and oats make a good response immediately after green manures, whereas wheat and barley usually do not respond so well, especially the first season after a green-manure crop. This brings us to a consideration of the effects of green manure on the yields of the succeeding crops.

**EFFECT OF GREEN MANURE ON YIELD OF CROPS**

Many experiments with various green-manure crops under a great variety of soil and climatic conditions have shown that, green manures, when properly used, increase the yields of succeeding crops and the effects are often noticed for several years. Under certain conditions, however, some crops immediately following legume green manures may be unfavorably affected. For example, when large quantities of highly nitrogenous material decompose rapidly in soil, the nitrate content may be increased to such an extent that crops do not grow normally; they frequently develop too luxuriantly in stems and leaves. The ripening of fruit may be delayed and the yield decreased; with small grains, lodging may result and the yield and quality of grain be diminished. The probable detrimental effect of turning under mature or carbonaceous materials has been discussed in a preceding section.

In regard to the effect of green manures on the yield of subsequent crops, two questions are pertinent.

**Questions**

1. How do green manures affect the yield of crops?
2. Do green manures have a pronounced residual effect on crop yields?

**Effect of Green Manures on Crops Immediately Following.**

The effect of green manures on the yield of subsequent crops has been studied by many of the state agricultural experiment stations, although a large share of the experimental work has been confined to the South and to the Atlantic seaboard. The work of nine southern states dealing with this subject has been summarized.<sup>1</sup> The increased yields resulting from turning under winter legume crops have ranged from 6 to 60 per cent over the yields of the check plots. In some of the southern states cotton yields have been increased from 22 to 100 per cent in various experiments by turning under legumes. The turning under of winter legumes has generally increased the yields of corn in the South from 24 to about 78 per cent. A summary of the effects of previous crops of summer and winter legumes on the yields of seed cotton and corn in the southern region and nearby states is presented in Table 30.

In the Corn Belt, where sweet clover is used extensively as a green-manure crop, substantial increases in corn yields usually are obtained. The turning under of summer legumes has also brought about marked

<sup>1</sup> "The Use of Cover and Green-Manure Crops," A. J. Pieters and R. McKee, *U.S.D.A. Yearbook* 1938, p. 436.

TABLE 30

EFFECTS OF PREVIOUS CROPS OF SUMMER OR WINTER LEGUMES ON THE YIELDS OF SEED COTTON AND CORN IN THE SOUTHERN REGION AND NEARBY STATES \*

| <i>Seed Cotton</i> |                     |                        |                     |                        |                               |
|--------------------|---------------------|------------------------|---------------------|------------------------|-------------------------------|
| Preceding Crop †   | Without Legumes     |                        | With Legumes        |                        |                               |
|                    | Crop acre, ‡ number | Yield per acre, pounds | Crop acre, ‡ number | Yield per acre, pounds | Difference in yield, per cent |
| Summer legumes     | 50                  | 756                    | 62                  | 1,163                  | +53.9                         |
| Winter legumes     | 106                 | 808                    | 247                 | 1,276                  | +58.0                         |

| <i>Corn</i>    |    |    |     |    |       |
|----------------|----|----|-----|----|-------|
| Summer legumes | 51 | 25 | 103 | 36 | +42.8 |
| Winter legumes | 53 | 25 | 175 | 35 | +41.2 |

\* Taken from "Summary of Effects of Legumes on Yields of Cotton and Corn in the Southern Region and Nearby States," Southern Region Agricultural Conservation, No. 3, 1937, p. 3, U.S.D.A., A.A.A.

† The crops of both summer and winter legumes were used in various ways—sometimes for hay, sometimes for seed; for many crops only the stubble was turned under, and for others the vines were turned under after the seed was harvested.

‡ Data in these columns represent the number of crops on an acre basis. This number was arrived at by the number of years the tests were conducted. For example, the number 50 in the table means that the yields of 50 crops of cotton, following legumes, from several experiments have been considered.

increase in crop yields, as has the turning under of the second growth of red clover. The great majority of records indicates that the turning under of a legume crop consistently increases subsequent crop yields.

Non-legume green-manure crops add organic matter to the soil but do not in themselves increase the nitrogen content of the soil above that existing at the time the green-manure crop was seeded. It is largely on this account that the increases in yields of the crops following non-legume green-manure crops are not so great as those of crops following legumes. As previously explained, the turning under of non-legumes frequently has resulted in a decrease in the following crop because of a

deficiency of nitrates. It is evident, therefore, that non-legume green-manure crops generally should be turned under when young or soluble nitrogen must be added to the soil to facilitate decomposition of the organic matter and to offset or prevent a nitrate deficiency.

**Residual Effect of Green Manures on Crop Yields.** In determining the returns from the practice of green-manuring, it is not sufficient to consider only the increase in the next crop. The green manure may have a marked effect on yields of subsequent crops for several seasons. The degree and the extent of the residual effect will be determined largely by the amount and kind of green manure turned under, the condition of the soil, and the climatic factors. Under otherwise similar conditions the residual effect probably would be extended over a longer period of years in cool humid regions than in warm humid regions.

The residual effect of green manures may be due in part to the improved mechanical condition of the soil or to any one factor or a combination of the other factors mentioned in the first part of this chapter (p. 261). At any rate, green-manuring is considered a part of a well-established system of soil management for maintaining or increasing the crop-producing ability of the soil.

# 12

## Farm Manures

Farm manure offers the farmer's chief opportunity for returning to the soil a portion of the plant nutrients removed in crops. Manure is one of the most important agricultural by-products; it is a perishable product and is frequently subjected to very severe losses. When calculated in terms of dollars and cents, the losses occurring with manure on American farms are astonishing. Only a fraction of the value of farm manure, perhaps not more than one-third, is actually realized. Farmers are acquainted with increased yields after an application of manure, but in general they do not understand the true nature of manure, the perishable nature of some of its constituents, or the losses occurring from its improper handling. In order to understand and appreciate better the value of manure, a few facts are presented regarding the production, losses, care, and field management of manure, from a practical point of view. The study may be pursued under the following divisions.

### Objectives

- A. Production of manure.
- B. Decomposition of manure.
- C. Losses occurring with manure.
- D. Methods of handling manure.
- E. Field management of manure.
- F. Fertilizing properties of manure.
- G. Effects of manure on the soil.

### PRODUCTION OF MANURE

It is common knowledge among farmers that soil fertility is maintained much more easily by a livestock system of farming than by a grain or cash system. This is true because a large share of the nitrogen and minerals in the crops fed and, to a less extent, of the organic matter is excreted by the animals. However, it is to be remembered that not all

the plant-food elements or the organic matter in the feed is excreted in the manure. Since the animal must be heated, have energy for performing work, and have nutrients for growth and body maintenance, it is obvious that a portion of the plant nutrients and organic matter is taken from the feed consumed by the animal. To obtain a better understanding of the real nature of manure it is necessary to consider the chemical changes certain constituents of feed undergo in the digestive tract of animals, the components of manure, the quantity and composition of excrements from different kinds of livestock, and the proportion of fertilizing constituents recovered from the feed eaten by the animals. Several questions present themselves in connection with a study of the production of manure.

### Questions

1. Which animals supply the largest proportion of farm manure?
2. Do pronounced chemical changes occur in the fertilizing constituents of feed in the digestive tract of animals?
3. What are the principal components of manure, and how do they differ in quantity and composition?
4. How do the quantity and composition of excrements from different kinds of livestock compare?
5. What percentages of the fertilizing constituents are recovered from feed consumed by animals?

**Quantity of Manure from Different Types of Animals.** The term farm manure has reference to the excrements from all animals of the farm. The total quantity produced in the United States from different kinds of livestock is in the following descending order: cattle, horses and mules, hogs, sheep, and chickens. Most of the manure that is eventually returned to the land is excreted by cattle and horses because, on the average farm, these animals consume a much greater portion of the grain and roughage and because the methods usually employed in handling such livestock make it easier to collect the manure. The quantity of manure excreted by 1,000 pounds live weight of different farm animals is given in Table 31 (p. 284).

**Chemical Changes in Feed in the Digestive Tract of Animals.** Complex chemical changes occur in the food in the digestive tract of animals. These changes are brought about partly by digestive enzymes and partly by the numerous bacteria which live in the intestinal tract. Twenty to 30 per cent of the dry weight of the solid excrement consists of living and dead cells of bacteria.

The various constituents of feeds undergo different rates and different degrees of decomposition. Sugars and starches are easily broken down; celluloses and hemicelluloses are less easily decomposed; and lignins are very resistant. Proteins vary considerably in their ease of breakdown. Proteins in "concentrates" are usually much more highly digestible than proteins in "roughages." Approximately one-half of the organic matter in the feed is decomposed during digestion.

Most of the potassium in feed is absorbed from the digestive tract and excreted in the urine. Only a small fraction of the phosphorus, except in hogs, is so absorbed. Consequently, most of the phosphorus in manure is carried in the solid fraction.

**Components of Manure.** Farm manure consists of two original components, the solid and the liquid. The solid material represents for the most part the undigested material, and the liquid portion represents the digested material that has been absorbed by the animal and then excreted.

The solid excrement, on the average, contains one-half or more of the nitrogen, about one-third of the potassium, and nearly all the phosphorus that is excreted by the animal. The nitrogen in the feces exists largely in two forms: first, the residual proteins that have resisted decomposition in the digestive processes; and, second, the proteins that have been synthesized in the cells of bacteria. Over one-half of the nitrogen may be present as synthesized protein, and this form is readily broken down when added to soils, so that the nitrogen is available to plants. However, the solid excrement contains large quantities of lignin, and it is believed that this lignin combines with protein, forming a lignin-protein complex which renders the protein less available. In other words, a large share of the organic matter in the feces is humified; a compound is formed similar to the humus that is formed in soils. As much as 50 per cent of the organic matter in the solid excrement may be in the humified state, and the nitrogen contained therein is only slowly available to plants when added to soils.

The liquid fraction or urine contains those plant nutrients which have been digested and utilized in the animal body and are later excreted. All the plant nutrients in this fraction are soluble and either are directly available to plants or readily become so. The liquid portion of manure differs from the solid not only in regard to the availability of nutrients but also in its low content of phosphorus and in its high content of potassium and nitrogen. (The distribution of plant nutrients between liquid and solid portions of manure is shown in Fig.

55.) In general, the more digestible the food consumed by an animal, the greater is the proportion of its plant nutrients that appears in the urine. Furthermore, as a rule, the higher the food is in nitrogen, the greater the digestibility of the nitrogen and the larger the amounts of nitrogen in the urine.

When voided, the nitrogen in urine exists largely as urea and hippuric and uric acids. These compounds are not volatile at ordinary temperatures, but manure contains organisms capable of rapidly breaking these compounds down with the formation of ammonia, which combines with water and carbon dioxide to make ammonium carbonate. This compound is unstable; even in solution it tends to decompose, losing ammonia, especially at higher temperatures [ $(\text{NH}_4)_2\text{CO}_3 \rightarrow 2\text{NH}_3 + \text{H}_2\text{O} + \text{CO}_2$ ]. This compound may lose all its ammonia on drying. The unstable nature of the nitrogen in urine presents a major problem in the handling of manure.

**Quantity and Composition of Excrements.** Many factors influence the quantity of manure voided and its composition, such as (1) the kind and age of the animal, (2) the kind and the amount of feed consumed, (3) the condition of the animal, and (4) the milk produced or the work performed by the animal. Wide variations are often found in the manure of animals even of a given class. Animals of different ages and doing different kinds of work require different amounts and proportions of nutrients to maintain them. A young animal, for example, which is building muscle and bone needs considerable phosphorus, nitrogen, calcium, and other elements, and the manure produced by such animals will contain much less of these elements. Since the composition of manure is so variable, data such as those presented in Table 31 can only be approximate.

The urine makes up only 20 per cent of the total weight of the excrement of horses but 40 per cent of that from hogs. These represent the two extremes. Since the urine makes up only 20 to 40 per cent of the total weight of manure from any animal, and yet contains approximately two-thirds of the total potash and somewhat less than one-half

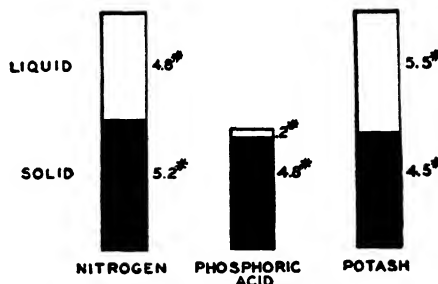


FIG. 55. Distribution of plant nutrients between liquid and solid portions of a ton of average farm manure.

## Farm Manures

TABLE 31

QUANTITY AND COMPOSITION OF FRESH MANURE EXCRETED BY VARIOUS KINDS OF FARM ANIMALS \*

| Animal  | Excrement | Pounds per Ton | Water, per cent | Nitrogen, pounds | P <sub>2</sub> O <sub>5</sub> , pounds | K <sub>2</sub> O, pounds | Tons Excreted † per Year |
|---------|-----------|----------------|-----------------|------------------|--|--------------------------|--------------------------|
| Horse   | Liquid    | 400            | ..              | 5.4              | Trace                                  | 6.0                      | ....                     |
|         | Solid     | 1,600          | ..              | 8.8              | 4.8                                    | 6.4                      | ....                     |
|         | Total     | 2,000          | 78              | 14.2             | 4.8                                    | 12.4                     | 9.0                      |
| Cow     | Liquid    | 600            | ..              | 4.8              | Trace                                  | 8.1                      | ....                     |
|         | Solid     | 1,400          | ..              | 4.9              | 2.8                                    | 1.4                      | ....                     |
|         | Total     | 2,000          | 86              | 9.7              | 2.8                                    | 9.5                      | 13.5                     |
| Swine   | Liquid    | 800            | ..              | 4.0              | 0.8                                    | 3.6                      | ....                     |
|         | Solid     | 1,200          | ..              | 3.6              | 6.0                                    | 4.8                      | ....                     |
|         | Total     | 2,000          | 87              | 7.6              | 6.8                                    | 8.4                      | 15.3                     |
| Sheep   | Liquid    | 660            | ..              | 9.9              | 0.3                                    | 13.8                     | ....                     |
|         | Solid     | 1,340          | ..              | 10.7             | 6.7                                    | 6.0                      | ....                     |
|         | Total     | 2,000          | 68              | 20.6             | 7.0                                    | 19.8                     | 6.3                      |
| Poultry | Total     | 2,000          | 55              | 20.0             | 16.0                                   | 8.0                      | 4.3                      |

\* Compiled from *Fertilizers and Crop Production*, Van Slyke, Orange Judd Publishing Co., 1932, pp. 218, 220.

† Clear manure without bedding; tons excreted by 1,000 pounds of live weight of various animals.

of the nitrogen, it is evident that, pound for pound, the urine is more concentrated and hence more valuable than the solid portion.

On a percentage basis there is considerably more nitrogen in the urine from most animals than in the feces, although more than one-half of the total quantity excreted is in the solid manure. The phosphorus, although existing in relatively small quantities, is found largely in the solid. Potash, however, occurs in larger quantity and in higher concentration in the liquid than in the solid.

It is interesting to note that water is one of the most variable constituents in manure. Poultry and sheep manure, on a tonnage basis, contain much greater quantities of plant food than any of the other manures; yet these two types of manure contain much less water. The different manures, on a wet basis, listed in Table 31, contain from 22

pounds of plant food per ton in cow manure to 47.4 pounds in sheep manure. If the composition of the total manure excreted by each of the different animals per 1,000 pounds live weight is calculated on a dry-weight basis, the manures would not vary so greatly in their plant-food content. Horse and sheep manures are considerably drier than cow or hog manures; this fact partially explains why horse and sheep manures heat more quickly than cow and hog manures in storage and why the former are preferred in the preparation of hotbeds.

**Fertilizing Constituents Recovered from Feed.** The fertilizing constituents in feed consumed by animals are not lost from the animal

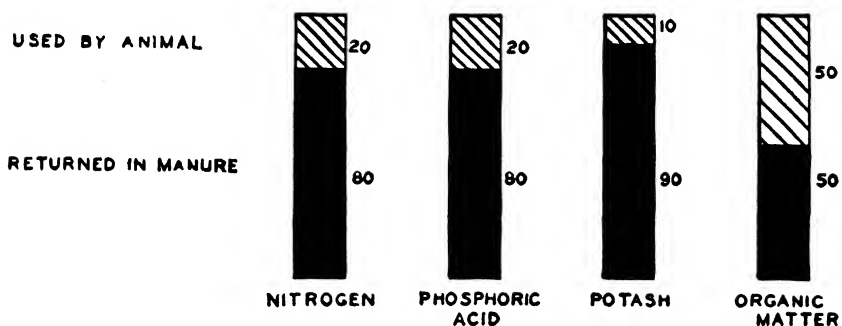


FIG. 56. Average proportion of plant nutrients and organic matter in feed consumed by animals which is excreted in manure.

body in any measurable quantity except through excretion in the urine and feces. Therefore, all the fertilizing constituents that are not built into animal tissue or secreted in milk are excreted in manure. The recovery of nutrients may be practically complete if the animal is not gaining or losing weight, producing milk, or supplying nutrients for developing young. Furthermore, animals that are not secreting milk and are losing weight may excrete more nitrogen than is contained in their feed as a result of animal-tissue breakdown. It has been indicated that young, growing animals and milking cows remove larger quantities of nutrients from feeds than do other classes of livestock. One thousand pounds of milk contain approximately 6.0 pounds of nitrogen, 1.7 pounds of phosphoric acid, and 1.7 pounds of potash; and about 2.5 pounds of nitrogen, 1.5 pounds of phosphoric acid, and 0.2 pounds of potash are contained in each 100-pound weight of a calf.

On an average, farm animals return in the manure from the feed consumed 75 to 80 per cent of the nitrogen, 80 per cent of the phosphoric acid, 85 to 90 per cent of the potash, and 40 to 50 per cent of the organic matter (Fig. 56). Variations in the recovery of nutrients

from feeds are due to different conditions, such as the kind and concentration of feed, age of animal, kind of animal, milk production, and work performed by the animal. Consequently, values may vary considerably from those given above.

### THE DECOMPOSITION OF MANURE

Many farm conditions do not permit the daily hauling of manure directly to the field; consequently, it must be stored, and, when handled in this way, marked changes occur in the manure. Owing to the heterogeneous nature of both the organic and inorganic constituents, the numerous kinds of microorganisms inhabiting manure, and the varying environmental conditions under which it is stored, a wide variety of chemical changes takes place. Some of the more important questions dealing with the decomposition of manure in storage may be listed as follows.

#### Questions

1. What changes are produced in nitrogen compounds?
2. What changes are produced in carbohydrates?
3. Do the mineral constituents undergo pronounced changes?
4. Does humus formation accompany the decomposition of manure?
5. What is the value of fresh compared to well-rotted manure?

**Changes Produced in Nitrogen Compounds.** The solid fraction of fresh manure as it comes from the stable consists of decayed or partially decayed plant materials with perhaps small quantities of broken-down animal tissue and bacterial cells. Usually this is mixed with litter and the entire mass is moistened with urine; it offers an excellent opportunity for many microorganisms to function. The rate and nature of changes affected by these organisms are determined to a large extent by the degree of aeration in the manure heap.

When manure is first produced, it is exposed to the air and rapid aerobic decomposition occurs with the liberation of considerable heat and carbon dioxide gas, and numerous changes in the nitrogen constituents. As the manure becomes more compacted, if it is kept moist oxygen is excluded and the decomposition becomes anaerobic. With the change from aerobic to anaerobic conditions, the rate of decomposition becomes slower, the temperature is lowered, and the nature of the decomposition products is materially different.

The urine fraction is relatively high in nitrogen, which exists chiefly in the form of urea. In decomposing, urea combines with water to

form ammonium carbonate, which in turn combines with carbonic acid to form ammonium bicarbonate. These carbonate compounds are unstable and tend to lose gaseous ammonia and carbon dioxide into the atmosphere. The odor of ammonia gas in horse stables indicates the volatile nature of these ammonium compounds. This tendency is more pronounced when temperature, alkalinity, and drying are increased. The decomposition of urea is materially reduced if the reaction is kept below pH 7.0. Under favorable conditions the change of urea to ammonium carbonate takes place very rapidly and may be almost complete within 3 or 4 days.

Organic nitrogen compounds other than urea occur chiefly in the undigested food protein and synthesized microbial protein of the solid excrement. They are both more or less resistant to biological decomposition, although some changes occur under both aerobic and anaerobic conditions. As decomposition proceeds, some of the nitrogen is released as ammonia and some is synthesized into microbial protein, with a net loss in total organic nitrogen. The foul-smelling compounds, skatole, indole, hydrogen sulfide, and amines are protein-decomposition products.

Nitrification in manure as a general rule is probably not an important process. A good supply of oxygen is essential for active nitrification; yet well-aerated manure heaps often develop high temperatures fatal to the nitrifying organisms. Therefore, nitrates are most likely to be produced in the outer parts of the heap where it is loose, reasonably dry, and relatively cool.

Nitrification presents the opportunity for denitrification, the formation by microorganisms of nitrogen gas from nitrites and nitrates. It has been suggested that, when nitrates are formed in the outer portions of the manure heap, they are transferred by leaching to the interior of the heap, where they are reduced to nitrogen gas under anaerobic conditions.

**Changes Produced in Carbohydrates.** The solid fraction of manure contains considerable quantities of cellulose, hemicellulose, lignin, and possibly lignin-protein complexes that escaped digestion and therefore represent the more resistant fractions of biological decomposition. Litters such as straw and corn stover, commonly used as bedding, contain relatively large amounts of celluloses and hemicelluloses and smaller amounts of sugars, starches, and proteins, all of which are rather easily decomposable.

The rate of decomposition and the nature of the products formed from carbohydrates depend mainly on the degree of aeration. Decom-

position is most rapid with a fairly high temperature, a good supply of air, ample moisture, and when the manure heap is loose and exposed to the open air. Under such conditions the decomposition of carbohydrates begins immediately unless delayed somewhat by a highly alkaline condition brought about by the decomposition of urea to ammonium carbonate. The principal change in the carbohydrates is induced by biological oxidation, the chief end products being carbon dioxide and water.

At first, when the more readily decomposable carbohydrates are being attacked, much energy is released in the form of heat, which may raise the temperature of the manure heap considerably. Since carbon dioxide, a volatile gas, is one of the chief carbohydrate-decomposition products, it is evident that as decomposition occurs there is a decrease in total dry matter. This decrease is rapid at first but slows up as the supply of readily decomposable carbohydrate is exhausted.

Anaerobic decomposition of carbohydrates is much slower than aerobic decomposition with the formation of different kinds of intermediate and end products. It is known as an acid type of fermentation, with the formation of carbon dioxide, methane, and hydrogen as end products and of various organic acids like acetic and butyric as intermediate products. These acids tend to combine with ammonia, lower the *pH* of the manure, and thus reduce ammonia losses. Anaerobic decomposition is accompanied by only slight increases in temperature and small losses of dry matter.

**Changes in Mineral Constituents.** Phosphorus is excreted largely in the feces, and it exists in both organic and inorganic combinations. During the biological decomposition of manure, organic phosphorus compounds are broken down with the formation of inorganic phosphates, which are usually considered more available. Decomposition affects the phosphorus compounds in manure mainly in a favorable way, and there is rarely, if ever, any danger of loss as the result of the decomposition processes.

A large part of the potassium of manure is voided in the urine, and most of the total present in mixed manure is soluble in water or at least is in a form which quickly becomes available for plant use. The potassium in the liquid portion usually is converted into potassium carbonate and may be active in neutralizing acids formed by the decomposition of carbohydrates. Insoluble potassium, calcium, and magnesium compounds, although occurring in only small quantities in manure, are changed during decomposition into more readily available forms. These compounds undergo no loss as the result of decomposition changes.

**Humus Formation in Manure.** During decomposition the fibrous portions of manure are disintegrated into a dark brown or black mass of fine, soft humus material, with the eventual disappearance of all traces of the original structures. This change is accomplished by a loss of carbon and hydrogen in the form of gaseous compounds. One-half of the original dry matter of manure may be lost in this manner. This explains the shrinkage in bulk of manure during decomposition.

The black colloidal organic substances in soils, commonly referred to as humus, are probably compounds essentially of lignin and protein, the latter being synthesized chiefly by microbial action. As has been mentioned (p. 282), one-half of the organic matter of feces may exist in a form similar to soil humus. Since considerable quantities of lignin are contained in both feces and bedding and since microbial protein may be synthesized in the rotting process, an increase in humus content can be expected to occur in manure.

**Value of Fresh vs. Rotted Manure.** In speaking of rotted manure, it is usually inferred that the original structure of the materials has disappeared more or less completely. In this connection, if we assume that the fresh manure is a normal mixture of urine and feces and that conditions have been controlled, fresh and rotted manure differ in composition as follows:

1. Rotted manure is richer in plant-food constituents. This concentration of plant nutrients is due to shrinkage in dry weight, which would automatically raise the level of plant food. One ton of fresh manure may lose one-half its weight in the rotting process. The loss occurs principally in the organic nitrogen-free constituents which contain no mineral plant-food elements.

2. More of the nitrogen in fresh manure is soluble. The decrease in soluble nitrogen is brought about by microorganisms in the synthesis of their body tissues during active decomposition of the organic constituents of the manure. There is considerable utilization of urinary nitrogen in the formation of complex proteins during the decomposition of manure. Unless rotting conditions are carefully controlled, there may be considerable loss in total nitrogen through volatilization.

3. The solubility of phosphorus is greater in decomposed manure. If no leaching occurs, there is no change in the total quantities of phosphorus or potassium.

That the increase in concentration of plant nutrients in rotted manure is obtained at the expense of large losses of organic matter and

considerable losses of nitrogen, especially available nitrogen, has not been fully appreciated. In general, it can be said that benefits derived by the rotting process may be more than offset by losses. If manure can be applied to the field daily, there is little justification for allowing it to rot except for special uses.

### LOSSES OCCURRING IN MANURE

Common methods of careless handling about the barn and delayed application of manure result in the dissipation of a large share of its value before it reaches the field. The degree to which manure decreases in value is determined mainly by the kind and by the methods used in collecting, storing, and applying it. Thus, several processes contribute to losses of the fertilizing constituents and organic matter in manure. Pertinent facts about these losses are brought out in the answers to several questions.

#### Questions

1. Do significant losses of the liquid portion of manure occur?
2. What losses occur by leaching?
3. Are volatilization losses important?
4. How can losses, due to scattering about barns and lots, be reduced?

**Losses of Liquid Manure.** The loss of urine from manure is serious from the standpoint of plant-nutrient content. This loss occurs mostly from failure to use ample bedding, leakage through stable floors, seepage into earth floors, or drainage from manure heaps.

On the basis of the total plant food in manure, about 50 per cent of the value is in the urine. It can be readily understood that large quantities of plant nutrients are lost and that the portions so lost, as already pointed out, are the most readily available forms of plant nutrients in manure. It is probably safe to assume that more than one-half the liquid is lost on many farms. In terms of dollars and cents, this represents a very significant loss to the farmer.

**Leaching Losses.** Frequently manure is thrown from stables into an adjoining lot, where it is unprotected from rain; sometimes it is thrown directly under the eaves of the barn. Leaching losses are greater when the manure is thrown in small, loose, and open piles. Very little excuse can be offered for storing manure in unprotected piles in the barnyard.

In the course of six months, manure which is exposed to rain may lose more than half its fertilizing value, depending on the quantity of

rain and the nature of the pile. The materials leached out are the most readily soluble and therefore the most quickly available plant nutrients. Leaching losses are confined not alone to the urine fraction but also to the nitrogen, phosphorus, and potassium compounds of the solid portion. Under alkaline conditions considerable quantities of soluble organic matter may leach out also; this is indicated by the dark color of leachings from manure piles.

**Losses by Volatilization.** Losses incurred by volatilization fall principally on nitrogen and organic matter. Large quantities of ammonia are produced in manure from urea and other nitrogenous compounds, and in the earlier stages of manure decomposition the ammonia is combined largely with carbonic acid as ammonium carbonate and bicarbonate. These ammonium compounds are rather unstable, and gaseous ammonia may be readily liberated. The tendency to lose ammonia nitrogen increases with the increase in concentration of ammonium carbonate and with the increase in temperature.

At ordinary temperatures little or no loss of ammonia from manure occurs at pH 7.0 or below. High temperatures, produced by aerobic decomposition in a loose manure pile, are conducive to very rapid loss of ammonia.

Freezing also tends to increase the loss of ammonia by increasing the concentration of the solution through the crystallization of the water. This loss may be considerable when manure is spread and becomes frozen.

Air movement greatly affects the loss of ammonia. Wind movements hasten evaporation of water, which decreases the capacity of the water to hold ammonia. Thus, manure which is permitted to dry out may lose appreciable quantities of ammonia. This fact emphasizes the importance of ammonia loss due to air circulation in loosely piled manure heaps and in overfermented manure that is forked frequently. Losses may also be considerable if manure is spread and permitted to dry before plowing under.

It has been pointed out (p. 287) that, when manure decomposes, it suffers important losses in organic matter and that these losses occur mainly in the carbohydrate constituents. One of the important end products of carbohydrate decomposition is carbon dioxide, most of which is lost from manure by volatilization. Shrinkage that accompanies the partial decomposition of manure is evidence of organic matter loss.

Little or no loss of phosphoric acid or potash from manure occurs through volatilization.

**Loss due to Scattering.** Another source of loss is that caused by the manure's being tramped into the mud of lots or dropped in roads or waste places where it cannot be picked up and returned to the land. Loss suffered by manure piled in lots may occur through the scattering of the material by chickens, hogs, and other livestock.

Some of this loss is unavoidable, but certainly part of it can be prevented. Such losses may be prevented or reduced by the direct hauling of the manure to the field or by protecting it from such mechanical scattering. Piling manure in straight-sided, compact, and flat-topped piles may also reduce such losses.

### METHODS OF HANDLING MANURE

The primary objective in the care and handling of farm manure is to prevent loss of plant nutrients as much as possible. Even under the most favorable conditions, it is practically impossible to prevent some loss of nitrogen and also some loss of organic matter. However, there is no difficulty in conserving all the phosphorus and potassium compounds because they are not volatile.

Although some loss of organic matter and nitrogen is inevitable under practical methods, such losses may be kept at a minimum with good management practices. Some of the more obvious ways of reducing losses of plant-food constituents in manure are indicated by the questions which follow. For the most part, these have been suggested by the discussion in the preceding section.

#### Questions

1. Does the use of litter decrease losses from manure?
2. Is it wise to haul manure directly from the stable to the field?
3. What are the proper methods for storing manure?
4. Are preservatives effective in decreasing plant nutrient losses?

**The Use of Litter.** Bedding or litter is used primarily to furnish clean and comfortable places for animals to lie down. In relation to the value of manure, bedding is used principally for these purposes: (1) to prevent loss of urine by drainage; (2) to make manure easier to handle; (3) to adsorb and hold plant nutrients; and (4) to increase organic matter and plant-nutrient content.

The principal value of litter as a preserving agent lies in its ability to absorb the urine, which in general carries more than one-half the fertilizing value of manure. The efficiency of various litters can be stated in terms of the amount of liquid which a given weight of mate-

rial will absorb. Ordinary straw can take up two or three times its weight of water; cut straw can take up about five times its weight of water; whereas good peat moss can absorb as much as ten times its weight of water.

The ability of litters to adsorb or fix both ammonia and potash protects them against loss by leaching, and the ammonia so fixed is less easily volatilized. This effect is very pronounced with such material as peat, but little fixation occurs with wood shavings, sawdust, corn stover, and straw.

Litter contributes some fertilizing constituents but usually in relatively small amounts, and the nutrients frequently are of low availability. Litter may also affect the process of biological decomposition of manures. The tendency for manure to heat, which increases ammonia loss by volatilization, and the change of inorganic to organic nitrogen are determined to a large extent by the readily decomposable material contained in the litter. Excesses of highly carbonaceous litters are to be avoided in order to prevent a decreased availability of nitrogen.

The minimum desirable amount of bedding to use is that quantity required to absorb all the liquid. This quantity depends mostly on the kind of litter used and the animal to be bedded. In general, straw, equal to 25 per cent the weight of excrement, will be sufficient to absorb the free liquid. The requirement of unchopped straw per head of livestock per day is: cattle, 9 pounds; horses, 10 to 12 pounds; sheep, 1 pound; and hogs, 1½ pounds. It is important that ample bedding be used; skimping in the use of bedding is false economy.

**Hauling Manure Directly to the Field.** Many farmers haul manure daily to the field, and this is usually a good practice. Manure spread on the land and worked into the soil is perhaps in its safest place. The soil has the capacity to fix rapidly large quantities of plant nutrients carried in the manure. As a rule very little loss occurs when manure is hauled directly to the field, except where it is spread on hillsides or on frozen ground, where it may be washed down the slope. If the liquid fraction has been thoroughly absorbed by bedding materials, there will be little loss by runoff. At any rate, it is better to have the liquid go into the soil for plant use than to let it soak in the soil of the barnyard or wash away in the drainage.

If manure is spread daily and is not worked into the soil, considerable loss of volatile ammonia may occur, owing to drying winds or freezing weather, or both, unless sufficient rain has fallen to wash the

ammonia into the soil. To decrease losses due to drying or freezing, the land should be disked after the manure is spread if possible.

Daily hauling of manure generally is regarded as the ideal method for preventing losses, although, as indicated above, there is considerable evidence that appreciable losses occur where spread manure remains unincorporated with the soil. However, since it is often impractical to apply manure daily during certain seasons of the year, it is necessary to resort to some storage on most farms.

If field conditions make it impossible to spread manure at certain times of the year, it can be hauled directly to the field and placed in flat-topped, straight-sided piles of 8 to 10 loads per pile. This method prevents scattering and washing and involves extra handling but usually is profitable.

**Storage of Manure.** In storing manure all practical precautions should be taken to keep losses at a minimum, and consideration must be given to costs involved. Handling manure not only increases the expense but also decreases the value of manure by exposing it to the air, thereby increasing decomposition losses.

Good storage of manure makes provision for keeping the manure heap (1) thoroughly compact, (2) sufficiently moist but not too wet, (3) under cover or shelter, and (4) undisturbed during storage.

Permitting the accumulation of manure under the feet of animals in stalls or in covered lots or pens is an efficient and practical means of storage. If plenty of bedding is used, the liquid manure is absorbed, the tramping of the animals compacts the manure, and fermentation losses are kept at a minimum. This type of storage also permits considerable flexibility in time of hauling. Some of the liquid manure may be lost when earth floors are used, and sometimes it may be economical to provide concrete floors to prevent this loss.

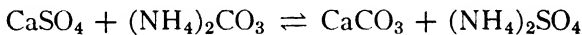
At least a temporary storage of manure is necessary on most farms. A common and wasteful method of storage is to place the manure in open piles in the barn lot or under the eaves of the barn. The principal menace of this type of storage is leaching. Storing manure under cover, in a closed shed, is the most practical method of preventing these losses. Such a shed should have a water-tight floor, four continuous walls, and a roof. The manure should be kept compact during storage.

On many farms it is necessary to resort to outdoor open storage. The manure should be placed in compact piles with straight sides and flat top and in such a position that it will receive neither roof nor surface water. Although this method usually results in considerable loss of plant food, loss is reduced if the manure is piled properly.

**Use of Preservatives.** Chemical preservatives are added to manure to decrease nitrogen losses. Their action may be due either to their prevention of the biological decomposition of urea and other nitrogenous compounds or to the conversion of volatile nitrogen compounds into non-volatile salts. To be most effective, these preservatives must be brought into contact with the liquid manure as soon as it is voided, because nitrogen losses may begin immediately.

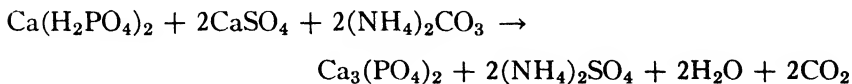
Strong acids such as phosphoric, sulfuric, and hydrochloric are effective preservatives. These chemicals make the manure acid, prevent urea decomposition, and change the ammonia compounds to non-volatile salts. Although these acids are effective preservatives, their cost and the difficulties encountered in handling them limit their practical use. It would appear, however, that sulfuric and phosphoric acids, particularly the latter, offer considerable possibilities. Phosphoric acid not only conserves nitrogen but also increases the plant-nutrient content of manure by supplying phosphorus, the element which is deficient in manure. The reversion of the phosphorus in phosphoric acid to an unavailable form does not accompany the fixation of ammonia.

Calcium salts of strong acids [CaSO<sub>4</sub>, CaCl<sub>2</sub>, and Ca(NO<sub>3</sub>)<sub>2</sub>] are of some value as preservatives. The reaction of CaSO<sub>4</sub> with (NH<sub>4</sub>)<sub>2</sub>CO<sub>3</sub> is



Their effect is that of changing the ammonia to a stable salt. As long as the manure is moist volatilization of ammonia will not occur, but if the manure becomes dry the chemical reaction is reversed and loss of ammonia may occur.

Superphosphate is used rather extensively as a preservative of manure. Its effect is similar to that of calcium sulfate (gypsum), although its action is more complex on account of the content of monocalcium phosphate in superphosphate. Ordinary grades of superphosphate (18 to 20 per cent) contain 40 to 50 per cent of gypsum, and they are more effective in preventing loss of ammonia, especially when manure becomes dry, than either gypsum or monocalcium phosphate used separately. The reaction of superphosphate with ammonium carbonate is:



In this reaction tricalcium phosphate is formed which does not react with ammonium sulfate on drying, so that loss of ammonia is prevented.

It is usually recommended that 1 to 2 pounds of 20 per cent superphosphate be used per day (24 hours) per horse or cow, and amounts for other animals in proportion to their weight. With horses the superphosphate is perhaps best applied just before the stables are cleaned; if the manure is allowed to accumulate in barns or lots, it is usually preferable to apply the phosphate just before each bedding. Dusting superphosphate in the gutter of the dairy stable after it has been cleaned is very satisfactory; it acts as a good absorbent of the liquid.

Peat has not only a high liquid absorption capacity but also a preservative action on manure. The peat may function as a preservative by absorbing ammonia nitrogen, by keeping the manure acid (in low-lime peats), or by virtue of the fact that it causes little heating when peat litter manures are stored. If bedding materials must be purchased, it is well to give peat adequate consideration.

### FIELD MANAGEMENT OF MANURE

Good management of manure does not stop at the barn or feeding pens; it extends to the field as well. The method of using manure in the field, with reference to crop, kind of soil, use of commercial fertilizers, and time, method, and rate of application, has much to do with getting the most benefit out of each ton. Questions most commonly raised in discussions of methods of handling manure are proposed here.

#### Questions

1. When should manure be applied, by what methods, and at what rate to give best results?
2. Should manure be supplemented with commercial fertilizers?
3. What crops respond best to farm manure?

**Application of Manure.** Prompt spreading of manure is generally considered best, although when in good storage it is likely to lose less value than if spread on the field without being plowed under or worked into the soil immediately. Losses of applied manure may occur in three ways: (1) volatilization of ammonia nitrogen as a result of drying or freezing; (2) surface runoff water carrying soluble portions of all three nutrients; or (3) leaching of nutrients.

**Time of Application.** Much experimental work with commercial fertilizers indicates that their effectiveness is decreased if they are applied a considerable time in advance of the seeding of the crop. This effect is attributed to leaching losses and to the fixation in less soluble forms of the plant nutrients by the soil. Fresh and properly stored manures contain large amounts of soluble nutrients, and the same prin-

ciples apply. Manure applied to corn land in the spring is likely to give greater returns than a like amount of manure applied in the fall. This effect will probably apply only to the crop of the first year and not to the later crops. If manure has already lost its readily soluble nutrients, the time of application is less important.

Little loss of nutrients from manure may be expected when applied on level, medium- to heavy-textured soils. These soils are safe for fall and winter spreading of manure, although they fix considerable quantities of the nutrients in less available forms. It is not advisable to place manure on sandy soils or on hilly fields much ahead of plowing time because of leaching and erosion losses. However, it must be remembered that manure applied on sloping fields decreases erosion losses, thereby counterbalancing to some extent the loss of nutrients from the manure.

*Method of Application.* In applying manure broadcast, it is important to get an even distribution over the field. As a rule, it is advisable to plow manure under or to disk it in immediately after spreading. The depth to which it should be worked into the soil depends on the climate, the type of soil, and the kind of manure. In general, manure should be incorporated in the soil so that sufficient air will be present for normal decomposition, and so that it is deep enough to prevent losses from drying. Deeper incorporations are more desirable on light, porous soils than on heavy or wet soils. The most important thing, in regard to the method and depth of mixing in the soil is the provision for completely covering the manure.

Although not substantiated by reliable evidence, it is commonly recommended that manure be thoroughly mixed with the surface soil; disk-ing before plowing is one method of attaining this end. There is some evidence to indicate that a higher efficiency can be obtained from localized placement (row or hill application) than from the broadcast application. Lack of necessary equipment for making such applications limits the use of this method at present, however.

A surface covering of manure, as a top-dressing on small grains in which a legume seeding is made, has been found an effective method of application in many areas. The improvement of moisture conditions and tilth of the topsoil brought about by the top-dressing more than compensates for the losses that may occur from the manure when this method is used. Although this is an effective method of improving stands of legume seedings in small grains, larger yields of small grains are often obtained from applying the manure before plowing than from top-dressing the crop.

*Rate of Application.* The rate at which manure should be spread will depend on the amount produced on the farm. As much of the cultivated land should be covered as possible. It is usually better to cover an entire field with a light application than to give only a part of the field a heavy coating. In other words, the crop response from 50 tons over 10 acres proves larger than from 50 tons on 5 acres. It is wise to gauge the rate of application so as to extend the acreage covered for each crop.

**Manure Supplemented with Commercial Fertilizers.** The content of phosphorus in manure is low compared with the amounts of nitrogen and potassium; for many soils and crops it is not a properly balanced fertilizer. For this reason it is usually advisable to supplement manure with a phosphatic fertilizer. The phosphorus may either be added with the manure as it is spread or applied to the succeeding crops at planting time. The practice of supplementing manure with phosphate is known as manure reinforcement. However, when the addition of nitrogen is the chief objective in the application of the manure, as when used in orchards or as a top-dressing on pastures, reinforcement may be unnecessary.

The conservation and wise use of farm manure alone will obviously not result in the maintenance of the plant nutrients in the soil. Since only a portion of the plant nutrients removed by crops is returned to the soil in manure, even when all the crops are fed on the farm, commercial fertilizers and lime, as needed, must accompany the use of manure if the proper level and balance of plant nutrients in the soil are to be maintained.

**Crops which Respond Best to Farm Manures.** Although all crops respond to manuring, the intertilled crops and grasses show most stimulation, and it is logical in the use and application of manure to favor those crops which are most responsive. Manure is essentially a nitrogen and potash fertilizer, and, since corn, cotton, tobacco, potatoes, and other commonly grown intertilled crops are heavy users of nitrogen and potash, it is obvious that these crops should be particularly benefited. When these crops are grown, they should receive at least a part of the manure that is produced on the farm.

Because of the tendency for oats to lodge when too much nitrogen is available, manure generally should not be used directly for this crop. Oats are seeded usually after some cultivated crop and feed quite satisfactorily on the residual manure which was applied for the cultivated crop.

Manure can generally be used to an advantage as a top-dressing on

fall wheat. It not only stimulates and protects the wheat but also greatly benefits the legume seeding that may be made in the wheat.

Manure is of value in orchards, especially on sandy soils where nitrogen and, sometimes, potassium give much concern. It may also supply some of the micro nutrient elements.

Where permanent pasture is a large enterprise, a liberal share of the manure can generally be used to advantage on the grass. Yearly dressings of finely divided manure, particularly if reinforced with phosphorus, have given excellent response.

Heavy applications of manure have proved desirable for truck, garden, and greenhouse crops. It is an excellent fertilizer material for intensive cropping operations.

### FERTILIZING PROPERTIES OF MANURE

Until the introduction of commercial fertilizers and the general utilization of leguminous crops, manure had always been the main source of supply of plant nutrients. In many places this is still true. The beneficial effects produced by manure on plant growth are believed to be due mostly to its content of nitrogen, phosphorus, and potassium. The growth-promoting influences of manure may be due in part to the presence of "minor" elements, organic matter, microorganisms, or to the presence of growth-regulating substances of a vitamin-like nature. As compared to ordinary grades of commercial fertilizer, manure is low in nutrient content. In fertilizer terms, it has on the average the analysis 0.5-0.25-0.5, containing  $1\frac{1}{4}$  units of plant nutrients as compared to about 20 to 30 units in ordinary mixed fertilizers. Although the nutrient content of manure is low, the quantity of nutrients added per acre in a normal rate of application of manure is usually five or more times the quantity normally applied as commercial fertilizer. This difference, however, is offset to some extent by the lower availability of some of the manure constituents. Several questions follow which may be of assistance in studying the fertilizer value of manure.

#### Questions

1. How does the availability of the plant nutrients in manure compare with that of the nutrients in commercial fertilizers?
2. What is the fertilizing value of manure compared to that of commercial fertilizers?
3. Are the growth-promoting effects of manure confined to its nitrogen, phosphorus, and potassium content?
4. Does manure have significant residual effects in soil?

**Availability of Fertilizing Constituents of Manure.** The nitrogen of the liquid fraction of manure is about equal in availability to that of mineral fertilizers like ammonium sulfate and sodium nitrate. The nitrogen of the solid fraction, however, is of very low availability. Experimental results reported show that less than 10 per cent of the nitrogen in the solid portion is recovered in the first crop, whereas over 70 per cent of the nitrogen from the urine, when applied with the solid, may be recovered in the first crop. As would be expected, the residual effect, measured by the recovery of nitrogen, is greater for the solid than for the liquid. The availability of nitrogen in ordinary farm manures may be expected to average about 25 or 30 per cent of that of mineral nitrogen fertilizers, depending on the kind of manure and the changes taking place in handling and storage.

The availability of the phosphoric acid and the potash of manure is essentially the same as that of mineral fertilizers. A number of investigators have reported a higher recovery by plants of phosphoric acid with manure than with chemical fertilizers.

**Fertilizing Value of Manure Compared to that of Commercial Fertilizers.** The value of a ton of farm manure in terms of dollars and cents has been the subject of much discussion. On account of the variable nature of manure, even though the amounts of plant nutrients present are known, it is difficult to place a value on the manure which expresses its worth to the farmer in the growing of crops.

Commercial values assigned manures usually are based solely on the quantities of plant nutrients contained and do not take into consideration the availability of the nutrients or the value of the organic matter present. Since the composition of stable manure is so variable, any estimate of value nearer than perhaps 50 per cent should be regarded largely as guesswork.

Crop increases per pound of nutrients applied frequently are somewhat higher with chemical fertilizers than with manure, perhaps owing to the unbalanced plant-nutrient ratio of manure for most crops and the lower availability of the nitrogen in manure. Obviously, the nature of the crop and soil conditions will affect any such comparison. The idea that applications of manures tend to increase yields more than commercial fertilizers in unfavorable seasons does not appear to be substantiated by experimental evidence. There have been no significant differences in the yield of corn and wheat on comparably manured and chemically fertilized plots in long-time experiments at the Ohio station.

**Growth-Promoting Effects of Manure Other than Those Attributed to Its Content of Nitrogen, Phosphorus, and Potassium.** Although crop increases per pound of nutrients applied are no larger and frequently are less with manure than with chemical fertilizers, it cannot be concluded that manure has no value other than its direct fertilizing effect. It would appear that the relatively low availability of a considerable portion of the nitrogen of manure should decrease, below the value usually found, the value of manure compared with chemicals. This may be offset partially by effects other than those produced by nitrogen, phosphorus, and potassium.

The benefits from *heavy* applications of manure on the physical properties of the soil frequently have been shown and generally are recognized by farmers. These effects, however, are not realized to any measurable degree under most farm conditions, especially on normally well aerated soils, because of the relatively small amounts of manure applied compared to the weight of the plow layer of soil.

It is believed that the addition of farm manures favorably influences the microbiological flora of the soil. Large numbers of microorganisms are introduced by manure, and this may hasten desirable biological changes in the soil. Certain organisms, thus added, convert organic nitrogen into ammonia; others change ammonia into nitrate nitrogen; and still others decompose carbohydrate materials. Manures also supply food materials for various desirable organisms already present in the soil. The increased production of carbon dioxide occurring in manured soils also may be of value in increasing crop growth.

The organic matter is an important factor in giving added value to manure in the soil aside from its direct effects on the physical and biological properties. When fresh manure is applied, the organic matter is gradually decomposed in the soil and its decomposition products tend to increase the availability of the mineral plant-food constituents in the soil. It has been demonstrated that the soluble humates in manure increase the solubility and mobility of mineral phosphates, and it has been suggested that the water-soluble and colloidal constituents of manure have a definite value aside from the accompanying nutrient salts.

There is also the probability that manure carries small quantities of certain growth-promoting substances, similar in nature to hormones and vitamins in animals. Thiamine chloride (vitamin B<sub>1</sub>), creatinine, and other growth-regulating substances are known to be present in manure in low concentrations. Whether these substances contribute anything of practical value to manure cannot be said now.

In addition to nitrogen, phosphorus, and potassium, manure contains calcium, magnesium, sulfur, iron, manganese, and other elements that no doubt have some fertilizing value. It is thus evident that manure may have value which cannot be attributed to its content of nitrogen, phosphorus, and potassium.

**Residual Effects of Manure.** An application of manure usually shows a favorable influence on crop yields for several years. These

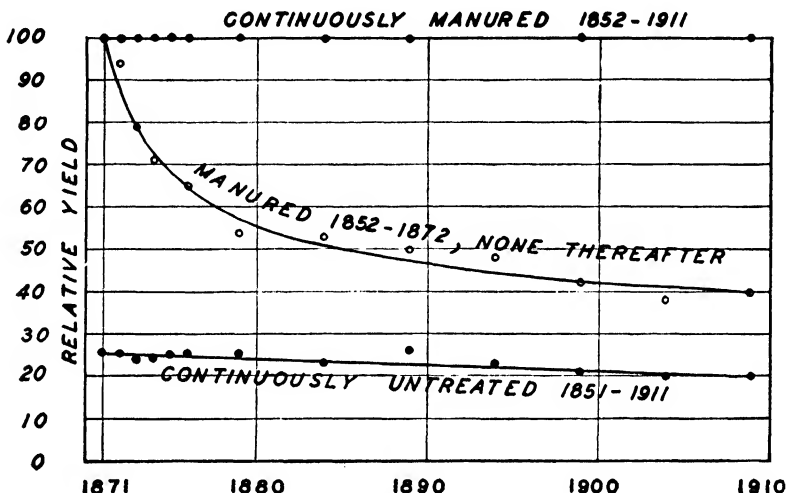


FIG. 57. Residual effects of heavy applications of manure (14 tons annually) on the yield of barley grown continuously on the soil of the Rothamsted Experiment Station in England. [From *Ohio Exp. Sta. Bull.* 605.]

beneficial effects are distributed over a longer time than those of chemical fertilizers. Striking results in showing the long-continued effects have been obtained (Fig. 57) by making heavy applications of manure for several years in succession and then discontinuing the application.

The lasting effect is due in part to the slow availability of certain plant-food constituents in manure, in part to the fact that a portion of the organic matter may remain for several years and aid in increasing the soluble plant nutrients in the soil, and in part to greater quantities of roots and stubble of the larger crops grown.

#### EFFECTS OF MANURE ON THE SOIL

Many chemical, physical, and biological properties of the soil are influenced by applications of manure. Perhaps the most important im-

mediate effects are chemical in that the quantities of available plant nutrients are increased. Immediate physical effects are less marked, although increases in aeration and the infiltration of water on heavy soils are observed. Increased crop growth, resulting from manure applications, tends to promote granulation and aids in retaining a good physical condition by protecting the soil from beating rains. The addition of manure also causes a rapid increase in the biological activity of the soil. The more important effects, however, are the result of long-repeated heavy applications of manure. The increased stable-humus supply of the soil serves as a storehouse for nitrogen and as an agent promoting desirable physical and biological properties. The following discussion of this topic should provide answers to the questions listed.

### Questions

1. Does manure increase soil organic matter and nitrogen?
2. Do applications of manure influence the mineral composition of soils?
3. Are the physical properties of the soil influenced greatly by the normal rates of applying manure?
4. How does the application of manure aid in decreasing soil erosion?
5. What is the role of farm manure in the maintenance of soil fertility?

**Effect on Soil Organic Matter and Nitrogen.** The value of organic matter in soils is generally recognized, and dark soils are usually considered superior to lighter-colored ones not so well supplied with organic matter. This fact, together with the observation that under most cropping conditions there is a gradual and continuous decrease in organic content, has resulted in placing much emphasis upon the humus-supplying power of manure. Available data indicate that this effect of manure has perhaps been exaggerated. The quantities of manure available in practice are normally too small to do more than retard the decline in soil organic matter. Therefore, it is more nearly correct to think of the normal applications of manure as a means of aiding in the maintenance of the soil organic matter supply rather than as a means of increasing it.

Changes in organic matter content of soils, attributable to manure, are affected in two ways: the actual residue of the manure applied, and the contribution by the roots and stubbles of the increased crops produced by the manure. The value of manure for conserving soil humus and nitrogen depends to a large extent on the crops used in the rotation. If manure is used on cultivated crops such as potatoes, corn, or tobacco, the gains from increases in roots and stubble residue may be very small,

but with such crops as the sod-forming legumes its maximum indirect effect is realized.

In regard to the effect of manure on soil nitrogen, it is to be kept in mind that only 75 or 80 per cent of the nitrogen of the food of animals can be found in fresh manure, and ordinarily a considerable portion is lost in handling. Although it is realized that raising crops on a farm and feeding them to animals of the same farm do not make so exhaustive a drain on soil fertility as removing them from the farm, yet under these conditions there can be no increase in soil nitrogen unless a large share of the acreage is devoted to leguminous crops. Even under a livestock system of farming there can be no effective increase in the soil nitrogen supply, unless much feed is purchased to be fed on the farm, a considerable acreage of leguminous crops is grown, or commercial nitrogen is purchased.

**Mineral Composition of the Soil as Affected by Application of Manure.** The extent of the changes produced in the mineral composition of the soil depends on the kind and amount of manure applied and on the accompanying losses of mineral elements from the soil. There can be no increase in the total mineral content of soils with a livestock system of farming where all the feed is grown and fed on the farm. In order that the total quantities of mineral elements be increased, it is necessary that they be purchased either in the fertilizer bag or in the form of feed to be fed on the farm. Obviously, the mineral content of a particular field may be increased by concentrating the applications of manure to that field, but that is merely taking fertility from one field and putting it on another.

Data from the Ohio station covering 32 years of cropping show that manure increased considerably the amounts of active calcium, magnesium, potassium, sodium, and manganese in the soil compared to the quantities contained in the unmanured soil. In most cases, however, even the heavier application (16 tons per five-year rotation) of manure did not maintain the supply of these elements at the original level. The lime requirement was not much affected by manure, but all plots were considerably more acid than at the beginning. The exchange capacity of the soil was increased.

**Effect of Manure on Physical Properties of Soil.** Some of the more important benefits from manure are brought about indirectly. The humus which manure supplies helps to improve or maintain the tilth or physical condition of the soil, the water-holding capacity, the permeability of the soil to water, aeration, and temperature relations. The physical properties of soils which are either too heavy or too light can

especially be improved by an increase in the humus content, which can be effected by large applications of manure.

With the usual limited quantities of manure available on general farms, marked changes in the physical properties of the soil are not realized. However, the physical effects of incorporating coarse organic matter with the soil are important even with moderate applications. Furthermore, manure, when used as a top-dressing, protects the soil from beating rains, decreases evaporation losses of water, and appreciably improves the tilth of heavy soils.

**Effect of Manure on Soil Erosion.** Incorporating manure with soil may be effective in reducing soil erosion by increasing the permeability of the soil to water, thus decreasing runoff losses, and by increasing the density of the vegetative cover, which in turn decreases rate of surface runoff and increases water penetration. A large number of experiments has been reported, showing that manure is effective in reducing both water and soil losses.

**Farm Manure and Maintenance of Soil Fertility.** As the average fertility of our soils has decreased, the saving and efficient use of manure have become a matter of vital importance in our system of agriculture. For many years manuring has been known to be perhaps the most logical and practicable method of aiding in the maintenance of soil productivity. Most cultivated soils give marked increases in yield where manure is applied, and this has brought about an increasing interest in the handling and care of this valuable product. Manure is no longer considered something to dispose of, but it is generally recognized as a most valuable by-product of the farm.

The greatest returns from manure are realized only when it is used in conjunction with other good soil-management practices. Proper methods employed in the handling, care, and use of farm manure are not sufficient in themselves to maintain soil fertility. Manure cannot be substituted for good cropping systems, lime, fertilizers, good seed, or proper drainage and cultural practices. Maximum returns from manure are assured only when it is well cared for, efficiently handled, and wisely used.

# 13

## Nutrient Requirement of Plants

The growth and development of plants are determined by numerous factors of soil and climate and by factors inherent in the plants themselves. Some of these factors are under the control of man, but many are not. Man has little control over air, light, and temperature, for example, but can influence the supply of plant nutrients in the soil. He may increase the supply of available nutrients by modifying soil conditions through good management or by making additions in the form of fertilizers. Anyone dealing directly with the growth of plants is particularly concerned with their nutrient requirements, and in this connection the following topics will be discussed.

### Objectives

- A. Elements used by plants.
- B. Effects of nitrogen, phosphorus, and potassium on plants and the quantities removed by crops.
- C. Determining soil-nutrient deficiencies.

### ELEMENTS USED BY PLANTS

If a soil is to produce crops successfully, it must have, among other things, an adequate supply of all the necessary nutrients which plants take from the soil. Not only must required nutrient elements be present in forms that plants can use, but also there should be at least a rough balance between them in accord with the amounts needed by plants. If any of these elements is lacking or if it is present in improper proportions, normal plant growth will not occur. Elements required by plants are designated *essential* or *indispensable*, and in giving consideration to them the following questions arise.

### Questions

1. From what sources do plants obtain their nutrients?
2. Which elements are essential?

3. Which of the essential elements receive major consideration?
4. Of what importance are the minor elements?

**Sources of Plant Nutrients.** Plants get their nutrient elements from three sources: air, water, and soil. Carbon and some of the oxygen are obtained from the air, whereas hydrogen, some oxygen, and possibly some carbon are taken from the soil solution. Legumes inoculated with effective nodule bacteria obtain some of their nitrogen from the air. Other nutrient elements must be taken from the soil by crops.

**Essential Plant-Nutrient Elements.** The ten chemical elements required in rather large quantities for growth of *all* crops are carbon, hydrogen, oxygen, phosphorus, potassium, nitrogen, sulfur, calcium, iron, and magnesium. Plant growth is limited by a deficient available supply of any one of these elements regardless of the abundance of the remaining ones. Experimental work before 1900 indicated that only these ten elements were required by plants, and they have been considered until rather recently the only ones necessary for the normal growth and development of plants. But, with the aid of improved technique, investigators have added to the original list several elements, including copper, boron, manganese, molybdenum, and zinc, which are undoubtedly essential although needed in small quantities. In addition to the elements listed which are required by all plants, certain others like sodium, chlorine, and possibly iodine are needed by some but not necessarily by all plants.

Although more than sixty elements have been detected in the analysis of plants, the mere presence of one of them in a plant does not prove that it aids in the development of that plant or that it is indispensable for plant growth. The necessity for any particular element can be determined only by carefully conducted experiments.

**Major Plant-Nutrient Elements.** Plants are made up principally of carbon, hydrogen, and oxygen, but some of the elements which occur only in relatively small amounts are just as essential to their growth and development as those which compose the greater part of the plant tissue. Obviously, it is incorrect to speak of one *essential element* as more important than any other *essential element* without specifying the viewpoint under consideration. From the farmer's standpoint, however, the elements nitrogen, phosphorus, and potassium are of major concern because years of experimental work and practical experience have proved that the available supplies of these elements in soils are more likely to be insufficient for maximum plant growth than are the supplies of the other essential elements. In fact, there are very few soils

which are not deficient in one or more of these elements. They assume agricultural importance because (1) they may be rapidly removed or lost from soils, (2) they may exist largely in unavailable forms in the soil, and (3) the only way to increase the phosphorus and potassium content of a soil is to buy them in some form or other. Consequently, the fertilizer companies have made these three elements (N, P, and K) the basis of the composition of commercial fertilizers which may supply any one, two, or all three of them. Calcium is also a mineral element greatly needed by many soils. Its importance is discussed on pages 137 to 141.

**Minor Elements.** It has already been mentioned that plants must have several elements other than nitrogen, phosphorus, and potassium for proper growth. In speaking of fertilizers, then, additional elements are referred to frequently as minor elements either because they are needed by plants in small quantities or because fewer soils are deficient in them, although in some soils the need for certain of them is as great as for the major nutrients. In fact, the known soil areas in which one or more of these elements limits crop production are increasing. However, undue publicity has undoubtedly exaggerated the general need for certain of these elements in fertilizer materials, and often the impression has been made in the minds of many people that soils in general are deficient in these elements.

Although minor or secondary elements such as magnesium, manganese, sulfur, calcium, zinc, copper, boron, iron, molybdenum, and perhaps others are essential, in many soils they are supplied abundantly by the materials commonly used in making mixed fertilizers or they are present in sufficient quantities in the soil. The addition of these elements to all fertilizer mixtures without regard to the crop or soil is unwarranted and sometimes may prove injurious.

Under conditions where a minor element deficiency occurs, two important questions come to our minds: (1) why are such deficiencies appearing now rather than at some earlier time; and (2) how do these elements act in promoting growth? There are perhaps several reasons why more minor element deficiencies are coming to attention now than at any previous time. The continued use of land for cultivated crops, the use of higher-analysis fertilizers (more pure salts), and the diminishing use of organic nitrogen fertilizers (plant or animal origin) have done much to hasten the occurrence of minor element deficiencies. It is also true that deficiencies occurred in some instances in earlier times, but the symptoms were not recognized.

The function of all these minor elements is not well understood, but

it is believed that they affect plant growth in one or perhaps more of the following ways: (1) are constituents of plant tissue; (2) act as catalysts or stimulants; (3) effect oxidation-reduction processes in the plant; (4) may aid in regulating the acid content of the plant; (5) may affect the plant osmotically; (6) may affect the entrance into the plant of other elements; or (7) may aid plant growth by providing a more favorable environment for the plant roots.

*Calcium* supply is frequently a limiting factor in acid soils. It is the important constituent of lime and is usually not classed as a fertilizer, but where the content of soluble calcium in soils is so low as to supply plants inadequately it should be considered a fertilizer. Calcium performs many functions in the soil and plant which are of great fundamental significance. It is generally present in large quantities in mixed fertilizers of the ordinary grades.

*Magnesium* is most likely to be deficient in sandy soils and particularly in seasons of heavy rainfall. Magnesium deficiencies have been observed frequently in the soils of the south Atlantic Coastal Plains and to some extent in the lighter soils in other parts of the country. This element is found in variable quantities in most fertilizers because of the frequent use of dolomitic limestone in their manufacture. Magnesium should be included in some form available to plants in nearly all fertilizers used on the soils along the Atlantic and Gulf coasts.



FIG. 58. In different soil areas animals may suffer from a deficiency of different nutrients in the locally grown feed. "A calf showing an advanced stage of salt sick or nutritional anemia. Note emaciation, lack of condition as indicated by the hair, appearance of the eye, and that the animal shows evidence of diarrhoea." The animal "recovered when given access to the iron-copper supplement." [Courtesy of Florida Experiment Station.]

*Manganese* usually is present in sufficient quantities in most soils, and it need not be added in fertilizer mixtures except in soils where a definite deficiency has been noted. The amount of manganese required by plants is small, and a deficiency in this element is most likely to be found in alkaline soils, especially those which originally were acid and have since been heavily limed. Manganese usually is needed for certain crops on alkaline muck soils.

*Copper, boron, and zinc* are sometimes found in insufficient quantities in soils, especially in the southeastern and in some of the eastern states. The yields of several crops grown on different soil types have been increased considerably by the addition of one or more of these three elements. The need for boron, especially for alfalfa growing, appears to be more widespread than it was originally thought. To soils in which deficiencies occur, these elements should be added in fertilizers in amounts found necessary by field tests.

*Sulfur* is used in large amounts by most plants but is usually found in considerable quantities in the soil. In areas close to industrial centers sufficient sulfur to supply crops is brought down by rain and snow from the atmosphere. The ordinary grades of superphosphate contain 40 to 50 per cent of calcium sulfate, and several other fertilizers and fertilizer ingredients also contain sulfur. They are all important sources of sulfur in the fertilizer-consuming areas of the United States. This element frequently is used as a corrective agent in overcoming an unfavorable soil condition known as alkali. As a result of the oxidation of applied sulfur to sulfuric acid, the pH of the soil is lowered.

*Iron* has been recognized as an essential element for a long time, and soils usually contain sufficient quantities of iron for normal plant growth. Its availability varies widely with the degree of soil aeration, being higher under anaerobic conditions. Occasionally soils are found which are deficient in available iron, and, on soils containing considerable quantities of lime, the movement and activity of the iron within plants are reduced in some manner by the presence of excess calcium. Investigations indicate a relationship between the solubility of iron and the supply of manganese. There is some reason for thinking that a deficiency of manganese in the soil leads to an iron toxicity.

#### EFFECTS OF NITROGEN, PHOSPHORUS, AND POTASSIUM ON PLANTS AND THE QUANTITIES REMOVED BY CROPS

In determining the kind and amount of fertilizer to use in any soil-management system, it is of interest and of practical value to have a

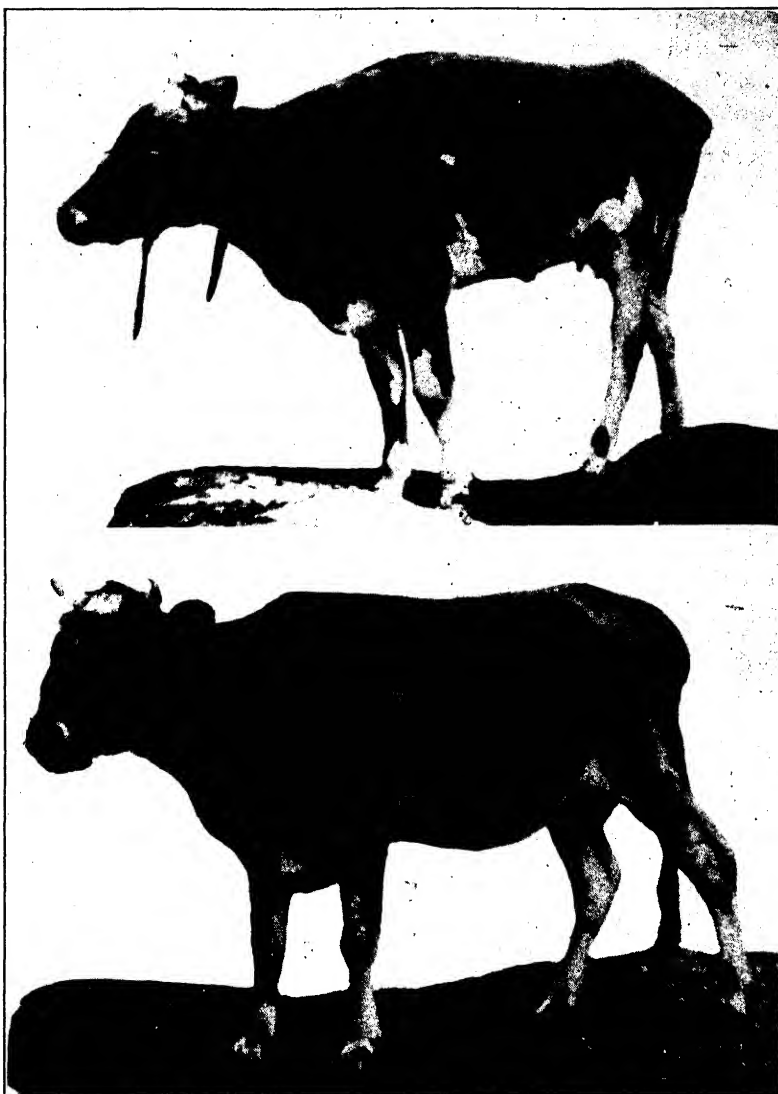


FIG. 59. Two views of the same cow. Above, when fed a ration deficient in cobalt. Below, after a cobalt supplement was included in the ration. [Courtesy of Dairy Department, Michigan State College.]

knowledge of the total quantities and of the proportion of nutrient elements removed from an acre of soil by good yields of crops. This factor must be considered in dealing with the problem of soil-productivity maintenance. A knowledge of the various ways in which the nutrient elements affect the growth of plants also is of practical interest. Each nutrient performs definite duties within the plant, and no one nutrient can be completely substituted for another. Although each element performs certain specific functions, all must work together to produce the best results. It must be remembered, therefore, that the effect of any particular nutrient on plant growth is governed by the supply of the other essential elements, and hence the effects of any one cannot be interpreted on the basis of the activity of that element alone. A balanced supply of nutrients tends to produce plants more resistant to diseases and attacks by insects. This section is concerned with the three elements (nitrogen, phosphorus, and potassium), and answers to the following questions will be sought.

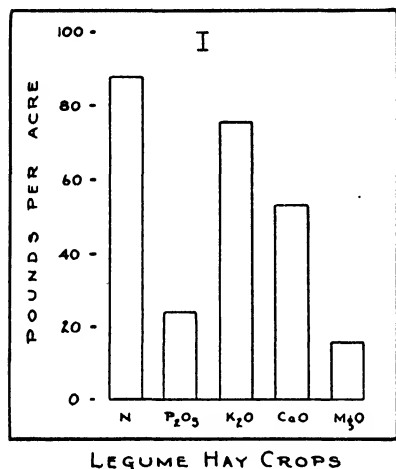
### Questions

1. Do crops remove appreciable quantities of nitrogen, phosphorus, and potassium from soils?
2. What are the effects of nitrogen on plant growth?
3. How do the influences of phosphorus differ from those of nitrogen?
4. Which roles does potassium play in plant growth?

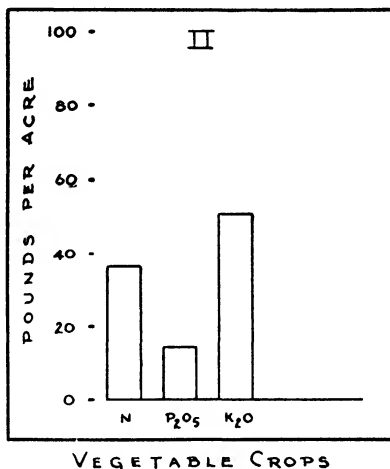
**Quantities of Nutrients in Crops.** Most of the plant nutrients removed from a virgin soil by the native forest or prairie vegetation are ultimately returned to the soil. Such soils in their natural, undisturbed condition are covered, at least in part, with a layer of organic matter in its various stages of decay, and, as decomposition proceeds, the nutrients in the organic matter are released and are again available for plant use. A decrease in soil fertility under virgin conditions is thus not likely to occur, owing to the return in time of almost all the vegetative growth.

When these soils are brought under cultivation, the condition is entirely changed because frequently most of the crop produced is removed from the soil permanently. Under a livestock system of farming, where the crop is fed on the farm, a considerable portion of the nutrients removed can be returned to the soil in the form of manure. But it should be remembered that the sale of any product off the farm, whether it is livestock, wool, milk, beans, wheat, cotton, tobacco, corn, sugar beets, vegetables, or fruits, represents a loss to the farm of the plant nutrients contained therein.

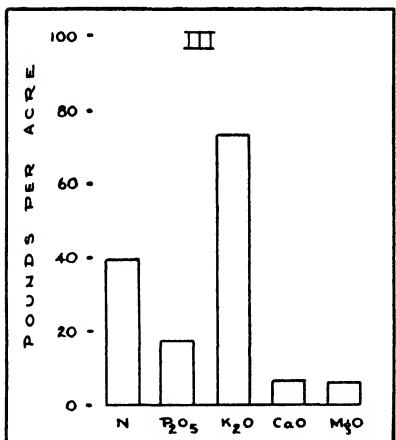
The nutrient content of several farm products is given in Table 32. In considering these quantities of nutrients, it is important to keep in mind that they are taken from the more readily available supply in the



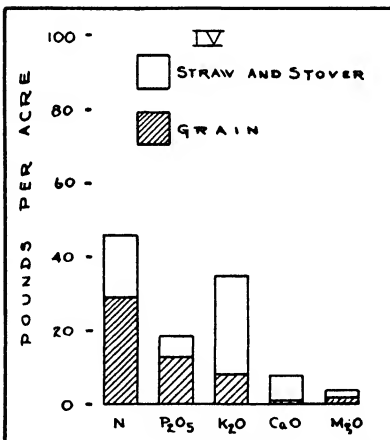
LEGUME HAY CROPS



VEGETABLE CROPS



TUBER AND ROOT CROPS



GRAIN CROPS

FIG. 60. The relative and also total quantities of plant nutrients contained in average yields of different crop groups vary greatly.

soil. Furthermore, the quantities removed by a single crop may seem rather small in some instances; but, when the quantities contained in all the crops of a rotation are totaled or when the amounts removed by crops for several years are considered, the necessity of supplying

## Nutrient Requirement of Plants

TABLE 32

## QUANTITIES OF NUTRIENT ELEMENTS CONTAINED IN FARM PRODUCTS \*

| Crop                                       | Amount     | Pounds               |   |                                      |                       |                         |
|--|------------|----------------------|---|--------------------------------------|-----------------------|-------------------------|
|  |            | Nitro-<br>gen<br>(N) | Phos-<br>phorus<br>(P <sub>2</sub> O <sub>5</sub> ) | Potas-<br>sium<br>(K <sub>2</sub> O) | Cal-<br>cium<br>(CaO) | Magne-<br>sium<br>(MgO) |
| Apple pomace, fresh (21.1% d.m.)           | 2,000 lb.  | 4.2                  | 6.4   | 2.0                                  | 0.4                   | ....                    |
| Apples                                     | 300 bu.    | 6.0                  | 3.0   | 15.0                                 | 1.2                   | 1.7                     |
| Barley, grain                              | 35 bu.     | 29.4                 | 12.6  | 8.4                                  | 0.7                   | 2.0                     |
| straw                                      | 1,600 lb.  | 13.4                 | 4.5   | 24.6                                 | 3.7                   | 1.1                     |
| Total                                      |            | 42.8                 | 17.1  | 33.0                                 | 4.4                   | 3.1                     |
| Barley, malt                               | 2,000 lb.  | 40.6                 | 8.4   | 7.4                                  | 1.2                   | ....                    |
| Beans, seed                                | 25 bu.     | 60.0                 | 18.0  | 19.5                                 | 3.0                   | 2.6                     |
| straw                                      | 2,000 lb.  | 28.0                 | 6.0   | 38.0                                 | ....                  | ....                    |
| Total                                      |            | 88.0                 | 24.0  | 57.5                                 | ....                  | ....                    |
| Beets, common, roots only                  | 25,000 lb. | 62.5                 | 25.0  | 125.0                                | 5.0                   | 5.0                     |
| mangles, roots only                        | 40,000 lb. | 60.0                 | 40.0  | 140.0                                | ....                  | ....                    |
| sugar, tops removed                        | 10 tons    | 36.6                 | 13.4  | 56.2                                 | ....                  | ....                    |
| Blackberries                               | 4,000 qt.  | 11.0                 | 3.0   | 12.0                                 | ....                  | ....                    |
| Bluegrass, Kentucky                        | 2 tons     | 53.0                 | 21.6  | 84.0                                 | 16.7                  | ....                    |
| Buckwheat, grain                           | 20 bu.     | 15.0                 | 6.0   | 3.0                                  | ....                  | ....                    |
| straw                                      | 5,000 lb.  | 62.5                 | 7.5   | 57.5                                 | 34.5                  | 6.0                     |
| Total                                      |            | 77.5                 | 13.5  | 60.5                                 | ....                  | ....                    |
| Cabbage heads                              | 20,000 lb. | 60.0                 | 20.0  | 80.0                                 | 18.0                  | 4.0                     |
| Carrots                                    | 10,000 lb. | 23.0                 | 13.0  | 53.0                                 | 6.0                   | 2.0                     |
| Cauliflower heads                          | 15,000 lb. | 42.0                 | 15.0  | 50.0                                 | 6.0                   | 3.0                     |
| Celery tops                                | 10,000 lb. | 25.0                 | 20.0  | 75.0                                 | ....                  | ....                    |
| Cherries                                   | 8,000 lb.  | 16.0                 | 4.0   | 20.0                                 | ....                  | ....                    |
| Corn, grain                                | 50 bu.     | 46.4                 | 18.2  | 11.0                                 | 0.4                   | 3.0                     |
| stover                                     | 3,000 lb.  | 30.0                 | 9.0   | 42.0                                 | 14.2                  | 2.5                     |
| cobs                                       | 500 lb.    | 2.0                  | 0.4   | 2.2                                  | 0.1                   | 0.1                     |
| Total                                      |            | 78.4                 | 27.6  | 55.2                                 | 14.7                  | 5.6                     |
| Cotton, lint                               | 500 lb.    | 1.0                  | 0.7   | 3.0                                  | 0.7                   | ....                    |
| seed                                       | 1,000 lb.  | 38.5                 | 14.8  | 12.0                                 | 2.8                   | ....                    |
| stalks and leaves                          | 2,000 lb.  | 28.5                 | 8.0   | 19.2                                 | 16.0                  | ....                    |
| Total                                      |            | 68.0                 | 23.5  | 34.2                                 | 19.5                  | ....                    |
| Cream (30%)                                | 1,000 lb.  | 3.9                  | 1.1   | 1.2                                  | 0.9                   | 0.1                     |
| Cucumbers, fruit                           | 100 bu.    | 5.5                  | 3.3   | 11.0                                 | ....                  | ....                    |
| Distillers' rye grains, dried (94.0% d.m.) | 2,000 lb.  | 58.0                 | 8.6   | 0.8                                  | ....                  | ....                    |
| Eggs                                       | 100 lb.    | 2.3                  | 0.4   | 0.15                                 | ....                  | ....                    |
| Fat, calf                                  | 200 lb.    | 4.9                  | 3.0   | 0.4                                  | 2.4                   | 0.1                     |
| lamb                                       | 80 lb.     | 1.6                  | 1.0   | 0.1                                  | 0.7                   | ....                    |
| pig  | 200 lb.    | 3.5                  | 1.3   | 0.3                                  | 0.9                   | ....                    |
| steer                                      | 1,000 lb.  | 23.3                 | 15.5  | 1.8                                  | 12.8                  | 0.4                     |
| Flax, grain                                | 15 bu.     | 30.5                 | 12.5  | 8.2                                  | 2.8                   | ....                    |
| straw                                      | 1,800 lb.  | 20.5                 | 3.4   | 18.8                                 | 13.2                  | ....                    |
| Total                                      |            | 51.0                 | 15.9  | 27.0                                 | 16.0                  | ....                    |
| Grape pomace, fresh                        | 2,000 lb.  | 19.0                 | 2.8   | 12.6                                 | ....                  | ....                    |
| Grapes                                     | 6,000 lb.  | 9.0                  | 6.0   | 18.0                                 | 0.4                   | 1.2                     |
| Hay, alfalfa                               | 3 tons     | 147.0                | 30.0  | 126.0                                | 83.5                  | 21.3                    |
| cowpea                                     | 2 tons     | 100.0                | 22.0  | 70.0                                 | 50.0                  | ....                    |
| lespedeza                                  | 2 tons     | 78.0                 | 41.0  | 81.6                                 | 55.6                  | ....                    |
| millet                                     | 3 tons     | 75.0                 | 27.0  | 90.0                                 | 23.0                  | ....                    |
| red clover                                 | 2 tons     | 84.0                 | 20.0  | 80.0                                 | 45.7                  | 10.8                    |
| soybean                                    | 2 tons     | 92.0                 | 28.0  | 44.0                                 | 49.2                  | 15.5                    |

TABLE 32 (*Continued*)

## QUANTITIES OF NUTRIENT ELEMENTS CONTAINED IN FARM PRODUCTS\*

| Crop                             | Amount    | Pounds               |   |                                      |                       |                         |
|----------------------------------|-----------|----------------------|---|--------------------------------------|-----------------------|-------------------------|
|                                  |           | Nitro-<br>gen<br>(N) | Phos-<br>phorus<br>(P <sub>2</sub> O <sub>5</sub> ) | Potas-<br>sium<br>(K <sub>2</sub> O) | Cal-<br>cium<br>(CaO) | Magne-<br>sium<br>(MgO) |
| Hay, sweet clover                | 2 tons    | 80.0                 | 22.0  | 74.0                                 | 31.4                  | 7.4                     |
| timothy                          | 2 tons    | 50.0                 | 22.0  | 40.0                                 | 7.1                   | 4.1                     |
| Leaves, scarlet oak              | 2,000 lb. | 31.4                 | 5.8   | 38.6                                 | 43.0                  | ...                     |
| red and black oak                | 2,000 lb. | 32.6                 | 5.0   | 39.8                                 | 46.2                  | ...                     |
| white oak                        | 2,000 lb. | 31.8                 | 3.6   | 29.2                                 | 53.4                  | ...                     |
| beech                            | 2,000 lb. | 35.8                 | 3.0   | 26.4                                 | 41.6                  | ...                     |
| sugar maple                      | 2,000 lb. | 28.0                 | 2.8   | 41.6                                 | 92.6                  | ...                     |
| red maple                        | 2,000 lb. | 11.2                 | 8.8   | 12.2                                 | 32.0                  | ...                     |
| Malt sprouts, dried (92.2% d.m.) | 2,000 lb. | 84.4                 | 14.2  | 30.4                                 | 4.8                   | ...                     |
| Milk                             | 1,000 lb. | 6.0                  | 1.7   | 1.7                                  | 1.1                   | 0.1                     |
| Muskmelons, fruit                | 5 tons    | 22.0                 | 8.0   | 40.0                                 | ...                   | ...                     |
| Needles, Jack pine               | 2,000 lb. | 11.6                 | 1.8   | 3.8                                  | 17.2                  | ...                     |
| Norway pine                      | 2,000 lb. | 13.4                 | 3.0   | 5.8                                  | 27.0                  | ...                     |
| white pine                       | 2,000 lb. | 12.8                 | 3.6   | 4.0                                  | 30.4                  | ...                     |
| Oats, grain                      | 50 bu.    | 32.0                 | 13.0  | 9.6                                  | 1.6                   | 1.9                     |
| straw                            | 2,500 lb. | 16.0                 | 5.0   | 31.2                                 | 7.5                   | 3.5                     |
| Total                            |           | 48.0                 | 18.0  | 40.8                                 | 9.1                   | 5.4                     |
| Onions, bulbs                    | 300 bu.   | 39.3                 | 15.4  | 37.6                                 | 18.8                  | 3.4                     |
| Parsnips                         | 6 tons    | 26.4                 | 24.0  | 78.0                                 | 9.6                   | 4.8                     |
| Peaches                          | 400 bu.   | 22.0                 | 11.0  | 45.5                                 | ...                   | ...                     |
| Pears                            | 300 bu.   | 7.5                  | 3.0   | 15.0                                 | 2.9                   | 1.4                     |
| Peas, canning, grain             | 20 bu.    | 44.0                 | 10.3  | 12.0                                 | 2.8                   | ...                     |
| straw                            | 3,000 lb. | 30.0                 | 5.7   | 31.8                                 | 59.0                  | ...                     |
| Total                            |           | 74.0                 | 16.0  | 43.8                                 | 61.8                  | ...                     |
| Peppermint                       | 2,000 lb. | 7.0                  | 4.0   | 13.0                                 | ...                   | ...                     |
| Plums                            | 200 bu.   | 15.3                 | 5.5   | 21.3                                 | ...                   | ...                     |
| Potatoes                         | 150 bu.   | 31.5                 | 13.5  | 45.0                                 | 1.8                   | 2.7                     |
| Raspberries                      | 4,000 qt. | 10.5                 | 4.5   | 12.0                                 | ...                   | ...                     |
| Rutabagas                        | 10 tons   | 40.0                 | 24.0  | 103.0                                | 12.0                  | 4.0                     |
| Rye, grain                       | 20 bu.    | 19.1                 | 9.8   | 6.7                                  | 0.5                   | 1.3                     |
| straw                            | 2,000 lb. | 10.0                 | 6.0   | 17.0                                 | 4.4                   | 1.4                     |
| Total                            |           | 29.1                 | 15.8  | 23.7                                 | 4.9                   | 2.7                     |
| Soybeans, grain                  | 15 bu.    | 79.5                 | 27.0  | 30.0                                 | 9.9                   | 12.6                    |
| straw                            | 3,000 lb. | 19.2                 | 3.9   | 18.6                                 | ...                   | ...                     |
| Total                            |           | 98.7                 | 30.9  | 48.6                                 | ...                   | ...                     |
| Strawberries, fruit only         | 6,000 qt. | 13.5                 | 10.3  | 27.0                                 | 1.4                   | ...                     |
| Sweet corn, ears                 | 4,000 lb. | 18.0                 | 8.0   | 12.0                                 | ...                   | ...                     |
| Tobacco, leaves                  | 1,500 lb. | 41.0                 | 6.8   | 72.0                                 | 57.0                  | ...                     |
| stalks                           | 1,250 lb. | 26.0                 | 5.7   | 39.6                                 | 9.7                   | ...                     |
| Total                            |           | 67.0                 | 12.5  | 111.6                                | 66.7                  | ...                     |
| Turnips, common                  | 10 tons   | 50.0                 | 20.0  | 90.0                                 | 10.0                  | 2.0                     |
| Wheat, grain                     | 25 bu.    | 30.0                 | 12.8  | 6.0                                  | 0.6                   | 2.0                     |
| straw                            | 2,500 lb. | 12.5                 | 3.8   | 15.0                                 | 5.2                   | 1.5                     |
| Total                            |           | 42.5                 | 16.6  | 21.0                                 | 5.8                   | 3.5                     |
| Wool, unwashed                   | 1,000 lb. | 54.0                 | 0.7   | 56.2                                 | 1.3                   | 0.2                     |

\* The data in this table are drawn or computed from data from many sources: *Fertilizers and Crop Production*, by Van Slyke; *Feeds and Feeding*, by Morrison; and other books and bulletins.

NOTE: d.m. means dry matter.

plant nutrients in the form of fertilizer and manure, to maintain soil productivity, is apparent.

The quantities of elements given in Table 32 do not represent the total quantities that crops require during growth, but rather the quantities contained in the harvested material. Roots and other portions of the plant which may not be harvested require considerable quantities of nutrients. Furthermore, with the advancing maturity of the plant, certain nutrient elements, such as potassium or calcium, may pass out of the plant roots into the soil.

Although a knowledge of the amount of plant food removed from a soil by the crops of any particular rotation will not tell definitely the fertilizer needs of the soil, this information, taken in conjunction with a knowledge of the plant-food condition in the soil, will give some indication of the amount of fertilizer and of the analyses to use. Obviously, the selling of plant food from the farm cannot go on indefinitely if soil productivity is to be maintained.

**Effects of Nitrogen.** Perhaps no element has received so much attention as has nitrogen in studies relative to plant nutrition. It is found in greater quantities in young, growing parts of plants than in the older tissues and is especially abundant in the leaves and seeds. Nitrogen is a constituent of every living cell, and hence its contribution to plant and animal life is evident. Its importance in crop production is emphasized by the knowledge that nitrogen generally occurs only in small quantities in soils in available forms, it is used by crops in large quantities, it is easily lost from soils by leaching and erosion, and of the three elements (nitrogen, phosphorus, and potassium) nitrogen is the most expensive to buy in the form of commercial fertilizers.

*Effect on Growth and Color.* A deficiency of nitrogen is evidenced by a gradual loss of chlorophyll which results in a light green to yellow color and by a slow and stunted growth. An abundance of nitrogen promotes rapid growth with a greater development of dark green leaves and stems. Although one of the most striking functions of nitrogen is the encouragement of aboveground vegetative growth, this growth cannot take place except in the presence of adequate quantities of available phosphorus, potassium, and other essential elements.

*Effect on Maturity.* An ample supply of available nitrogen during the early life of the plant may stimulate growth and result in earlier maturity. However, the presence of an excess of nitrogen throughout the growing season of the plant frequently prolongs the growth period, and the plant may fail to mature properly. This effect is especially significant for certain crops in regions having a short growing season.

or in areas where an early-fall freeze may do great damage to fruit trees whose season's growth period has been prolonged.

*Effect on Quality and Disease Resistance.* A large supply of available nitrogen encourages the production of soft, succulent tissue which is susceptible to mechanical injury and the attack of disease. Either effect may decrease the quality of the crop. However, the development of softness in the tissues may be desirable or undesirable, depending on the kind of crop. For vegetables used for their leaves, pronounced succulence, tenderness, and crispness are desired. Other vegetables and some fruits may have their keeping and shipping qualities impaired when they are grown with an excess of available nitrogen. An excess of nitrogen may encourage lodging in grains, which frequently decreases the quality, but a normal amount of nitrogen usually increases plumpness in grains.

**Effects of Phosphorus.** The total supply of phosphorus in soils is relatively small, and the available supply usually falls short of crop requirements. Phosphorus is used more generally on mineral soils than is any other fertilizing element, and it is usually the first element that becomes deficient after a soil is brought under cultivation. The importance of phosphorus in plant and animal nutrition is perhaps well recognized by every one. It is present in seeds in larger amounts than in any other part of plants, although it is found extensively in the young growing parts. Like nitrogen, it is a constituent of every living cell.

Sufficient quantities of available phosphorus are necessary for normal transformations of carbohydrates in plants—the changing of starches to sugars, for example. Phosphorus is necessary also for the assimilation of fats in plants, and apparently it increases the efficiency of the chloroplastic mechanisms. Phosphorus enters into the composition of nuclear compounds in cells. The presence of phosphorus aids the plant in taking up potassium and tends to counteract the effects of excess nitrogen. An excess of phosphorus in proportion to the supplies of other required nutrients may decrease yields, especially on the lighter soils. This is believed to be caused by a hastening of the maturation processes and by the consequent reduction in vegetative growth. All the functions of phosphorus in plant nutrition and their relationships are not well understood. The effects of phosphorus on plants are less striking than those of nitrogen and are often difficult to recognize by casual observation.

*Effect on Root Development.* The most obvious effect of phosphorus is usually on the root system of plants. Phosphorus-starved plants tend to have a stunted root system which decreases their feeding zone, and

thus the plants are less able to withstand adverse conditions. The stimulation of root development, due to the addition of available phosphorus is especially valuable in the heavier soils where root development is naturally restricted.

*Effect on Growth.* Phosphorus hastens the ripening processes of plants. In the presence of sufficient available phosphorus, seed formation begins sooner and crops may mature several days earlier than where phosphorus is deficient. Phosphorus is essential for seed formation, and its effect on the maturity of crops may be explained on this basis. In certain climatic regions the importance of hastening maturity is obvious. Since phosphorus is essential for cell division, plants starving for this element are stunted in growth.

*Effect on Quality and Composition.* Phosphorus-starved plants may mature late and be injured by frost; thus the quality is lowered. Hay and other forage crops grown on phosphorus-deficient soils contain reduced amounts of this element and so are of inferior feeding value. The presence of this element seems to favor good pollination, which affects quality and yield in corn. It strengthens the straw and stems and thus decreases lodging, decreases the ratio of straw to grain in cereals, and increases total yield. In general, phosphorus improves the quality of plants and plant products by stimulating the production of a more vigorous plant growth, making it more disease resistant.

**Effects of Potassium.** The total content of potassium in soils is usually high, but because of the small amounts available to plants it is of major concern to farmers. Potassium is usually needed more often and in larger quantities in the light soils (sandy soils) and mucks than in the heavy soils. The total potassium in sandy soils frequently is low, and that of muck often reaches a very low level. Potassium, in  $K_2O$  (potash), makes up about 40 per cent of the ash of most plants, and it is not localized in any part of the plant to the extent that phosphorus is, although in some crops it may tend to accumulate in the leaf and stem rather than in the grain.

With the passing of time the need for potassium in our cultivated soils is becoming more acute. Potash-deficiency symptoms in plants are appearing in widespread areas. This element plays an important part in many of the vital physiological processes in the plant, although the exact nature of the mechanism by which potassium functions is not definitely known. It is essential in all cell metabolic processes and apparently has a specific role in influencing the uptake of certain other mineral elements, in regulating the rate of respiration, in affecting the

rate of transpiration, and perhaps also in influencing the action of enzymes.

Potassium encourages the development of the root system of plants. An excess of this element may delay maturity, although in general it has a balancing effect on both nitrogen and phosphorus with respect to the maturation processes. The intake and retention of water seem to be regulated to some extent by the quantity of potassium present; thus it affects the resistance of the plant to injury from drought and frost.

*Effect on Synthetic Activities.* One of the most important functions usually attributed to the element potassium is its effect on the plant synthesis of carbohydrates and proteins. Potassium is essential for the production of starch, sugar, and other carbohydrates and in the translocation of starch and other materials within the plant. Some investigators are of the opinion that potassium aids in the reduction of nitrates in the plant preparatory to protein synthesis. It is also believed essential in the development of chlorophyll and in the synthesis of oils or fats and albuminoids. Potassium is not found in permanent organic combinations with these compounds, but it is believed essential for the production of them all.

*Effect on Vigor and Disease Resistance.* Potassium appears to improve the general tone and vigor of the plant, which in turn permits the plant to be more disease-resistant. As a rule, crops which do not receive sufficient potassium are more susceptible to disease. This effect is especially noticeable where the plant receives excess nitrogen.

*Effect on Quality and Composition.* Potassium increases plumpness in grains, producing greater test weight per bushel, and it makes the stalk and straw of plants more rigid, thus preventing lodging to a certain extent. Ears of corn produced on potassium-deficient soils frequently are chaffy, tapering at the tips, and the kernels are loose on the cob because they are not well filled with starch.

Where a deficiency exists, the addition of potassium affects the quality of tobacco, improves the quality in potatoes, and increases the sugar content of sugar beets. Many other illustrations might be given to indicate the effect of potassium on quality and composition of crops.

#### DETERMINING SOIL-NUTRIENT DEFICIENCIES

Since the growth of crops is influenced by rainfall, drainage, temperature, sunlight, rotation, soil reaction, previous soil treatment, and bacterial activity, in addition to the supply of nutrients, it is a very diffi-

cult matter accurately to determine the soil conditions that require the application of fertilizers. The practical problem in regard to the conditions that require fertilizer application is to know beforehand or at planting time the kind and quantity of fertilizer to use to give the most satisfactory yields. In working toward that goal, due consideration must be given to the crops and to the soil; they are of prime concern. Even after these things are taken into consideration, the results obtained from fertilizer applications are frequently affected and to a large extent determined by climatic conditions.

Not one of the several laboratory methods that have been developed is able to answer, on the basis of chemical or biological principles, the practical question of just what fertilizer to apply to a given soil to get the maximum yield of a certain crop under field conditions. This should not be expected, for the supply of nutrients is only one of the many factors that affect plant growth. The fact remains, however, that certain chemical tests aid in giving us a better understanding of the possible supply of nutrients and permit the determination of the presence of certain harmful substances. Some questions which come up in connection with the determination of nutrient deficiencies in soils are suggested.

### Questions

1. Of what value is a total chemical analysis of a soil?
2. Do the rapid chemical tests aid in determining soil deficiencies?
3. What are the limitations in determining these deficiencies by analyzing the ash of plants?
4. What are the possibilities and limitations of testing the sap of plants and plant tissue?
5. Can plant symptoms be used as a guide in diagnosing soil-nutrient needs?
6. What objections may be raised to the use of pot tests?
7. Why are field tests regarded as the most reliable method?

**Total Chemical Analysis of Soil.** The plant-nutrient content of a soil is usually divided into two arbitrary groups: the potential supply and the available supply. The potential supply is determined by a total chemical analysis of the soil and tells us very little about the fertilizer needs of the soil. It has already been emphasized that the growth of plants is determined by the nature of the chemical changes taking place in the soil and plant, and not by the total or potential plant-nutrient content of the soil. A total chemical analysis of a soil tells us little about the rate at which plant nutrients will become avail-

able during the growing season, although, if good growing conditions prevail, a soil analyzing high in plant nutrients would generally be expected to produce more available nutrients during the growing season than a soil low in total plant nutrients. However, a total chemical analysis of soils is of value to the soil scientist and hence serves a useful purpose although it is of little service to the farmer.

**Rapid Tests.** Various methods have been developed to measure the so-called available<sup>1</sup> plant-nutrient supply in the soil, and attempts have been made to interpret the test results in terms of the fertilizer requirements of the soil. To diagnose soil conditions by means of chemical methods is to attempt to place the matter on a basis which may apply to all soils. Although these tests are of value in determining the chemical condition of nutrients in a soil at any particular time, none of them can be relied on entirely as a basis for making fertilizer recommendations. It is impossible, by any known chemical or biological soil test, to imitate the action of higher plants in obtaining nutrients from the soil and so to determine the available plant-nutrient supply. It is noteworthy, however, that much progress has been made in the practical interpretation of soil-test results.

Many systems of rapid chemical tests have recently been developed for determining nutrient deficiencies, nutrient levels, and the presence or absence of harmful substances in soils, the last being particularly important from the standpoint of greenhouse soils. The main difference in these various methods lies in the strength and nature of the extracting reagent used. Such tests should be looked on as guides in determining fertilizer needs but should not be relied on entirely. The proper interpretation of results obtained by using these rapid methods is often rather difficult and should be attempted only by experienced persons.

**Testing the Ash of Plants.** It is generally understood among investigators that chemical analyses of plant ash cannot be depended on to indicate the need for fertilizers. The proportion of various elements that may be found in plants depends on numerous and varied factors,

<sup>1</sup> Nutrients that are in a condition suitable for immediate use by plants, or those in a state that can be readily changed under good soil conditions to suitable forms for plant use, are commonly called *available*. Those compounds or materials which are not in form for immediate use and are not readily changed to usable forms are called *unavailable*. The line of division between availability and unavailability is quite arbitrary, to say the least, and it is impossible always to distinguish between the two since so many factors must be taken into consideration.

such as (1) general climatic conditions, (2) species of plant, (3) nature of root system, (4) stage of maturity of plant, (5) methods of cultivation, and (6) nature of soil. In spite of the fact that analyses of plant ash cannot be used in general as a basis for fertilizer recommendations, they may be used as an aid in explaining unproductive areas in certain localities. The relation of phosphate content of feeds to the phosphate content of soils, on the one hand, and the health and nutrition of animals, on the other, may be given as an example.

**Sap and Tissue Tests.** Methods involving the testing of the expressed sap of plants and of plant tissue have been developed with the idea of aiding in making fertilizer recommendations. Obviously, several limitations have presented themselves in these attempts. These tests are of value in that they indicate the content of elements in soluble mineral forms in the plant, and before an element can be of direct benefit it must be taken up by the plant. In this respect they are superior to soil tests, which do not tell the amounts of nutrients a plant will absorb. However, tests of plant tissue do not permit interpretation at seeding time, which is important from the standpoint of determining the immediate fertilizer requirements. These test results may indicate certain limiting elements which may be used as a guide in making fertilizer recommendations the following year on the same crop and may perhaps be of value in indicating the kind of fertilizer to apply as a top-dressing or side dressing the current season.

In regard to tests on the sap and tissue of the plant, the matter of determining the part of the plant to test and the stage of maturity of the plant when samples should be taken must be considered. Regardless of the various limitations of these rapid plant tests and also of the rapid soil tests, when the two are used simultaneously they offer much information regarding the chemical condition of the soil and are most valuable guides in making fertilizer recommendations. Plant-tissue tests are also of great value in substantiating diagnoses of deficiencies and toxicities based on foliar symptoms.

**Deficiency Symptoms.** When plants are starving for any particular nutrient (nutrients are discussed elsewhere), characteristic symptoms usually appear on these plants. If crops are not vigorous and healthy, it is important to know and understand the cause. If the unhealthy appearance is due to disease, it may be possible to save the crop by spraying, or, if it is a nutrient deficiency, fertilizers may be applied as a top-dressing or a side dressing in time to save the crop. These deficiency symptoms appear only when the supply of a particular element is so low that the plant can no longer function normally, and then it will



FIG. 61. Bean leaves yellowed from deficiency of manganese and potassium compared with a normal leaf on the left. Center leaf yellowed first at the tips and along the edges. The crinkled appearance was caused by the continued growth of the interior portions after growth had ceased along the leaflet edges. The right leaf is uniformly yellow and has green veins due to insufficient manganese. [*Michigan Spec. Bull. 353.*]

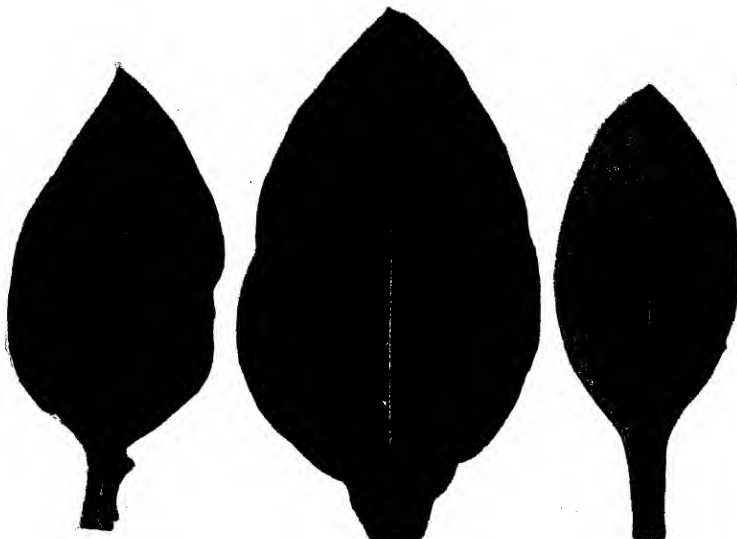


FIG. 62. Tobacco leaves indicating deficiency of iron and manganese. Center leaf is normal. The left leaf has lost its color except along the principal veins because of insufficient iron. The checkered appearance and associated dead spots in the right leaf are due to manganese deficiency. [From *Hunger Signs in Crops*, Kodachrome by J. E. McMurtrey.]

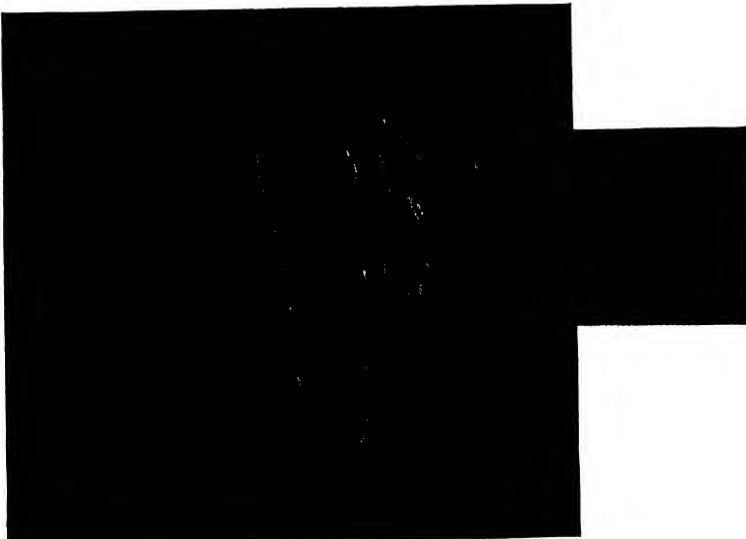


FIG. 63. Corn leaves deficient in nitrogen, potassium, and phosphorus. The left leaf is normal. The yellowed tip and margins of the center leaf arise from lack of potassium. In the next leaf insufficient nitrogen caused yellowing to affect the tip and then proceed up the midrib. The insert on the right shows the reddish purple edges of a phosphorus-deficient leaf next to a normal leaf. [*Michigan Spec. Bull. 353.*]

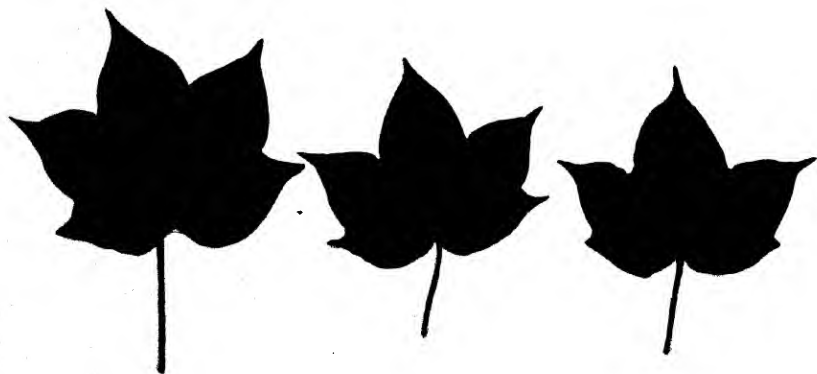


FIG. 64. The cotton leaf on the left is normal. The other two show a purplish red color with green veins because of insufficient magnesium. [From *Hunger Signs in Crops*, Kodachrome by J. E. McMurtrey.]



FIG. 65. (*Left*) Alfalfa and clover leaves showing potassium deficiency. The silt loam soil was fertilized with phosphate. The white dots scattered through the yellow edges of the leaflets are unmistakable evidence of potassium shortage.

FIG. 66. (*Right*) Wheat grown on clay loam soil. The yellow, dried-up lower leaves of the left bunch are characteristic of nitrogen deficiency. Tissue tests showed a high nitrate content in the bunch of plants to the right and no nitrate in those of the left bunch. [*Michigan Spec. Bull. 353*.]

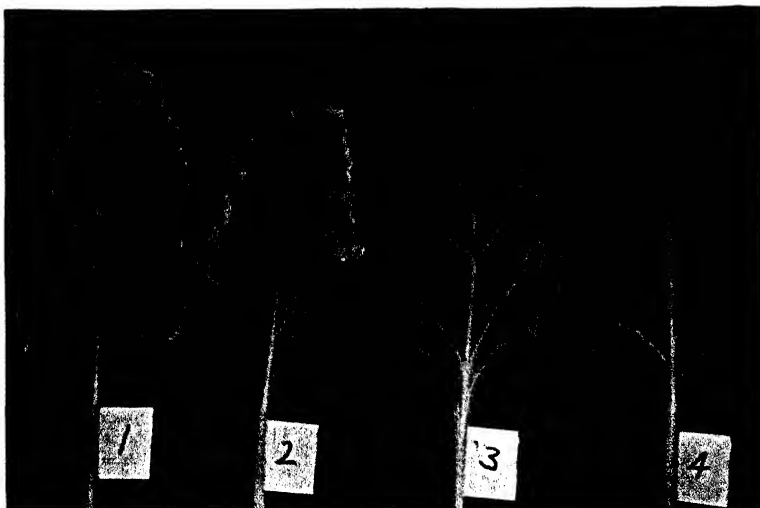


FIG. 67. Sugar beet leaves deficient in (1) manganese, (2) potassium, and (3) nitrogen. Leaf number 4 is normal. Note that when nitrogen is deficient the veins do not remain green as they do when manganese is deficient. The marginal yellowing of the potassium-deficient leaf is characteristic. [*Michigan Spec. Bull. 353*.]

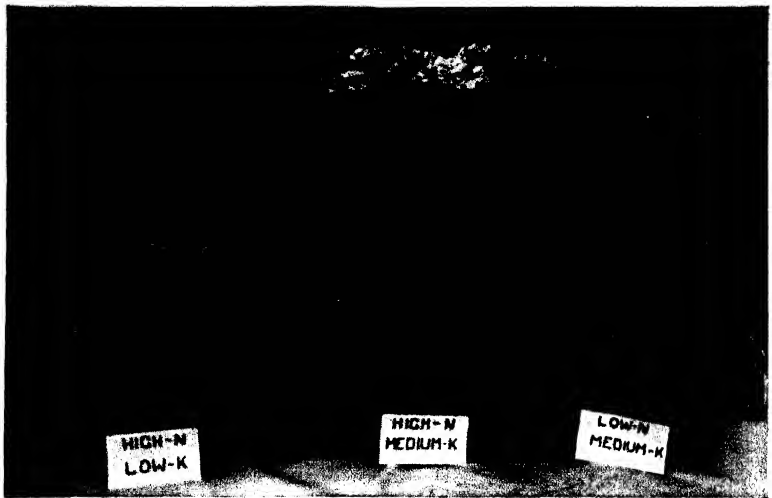


FIG. 68. Cineraria plants, potassium-deficient on the left, normal in the center, and nitrogen-deficient on the right. A normal color in the foliage of flowering plants is important from a sales standpoint. [*Michigan Spec. Bull. 353*.]

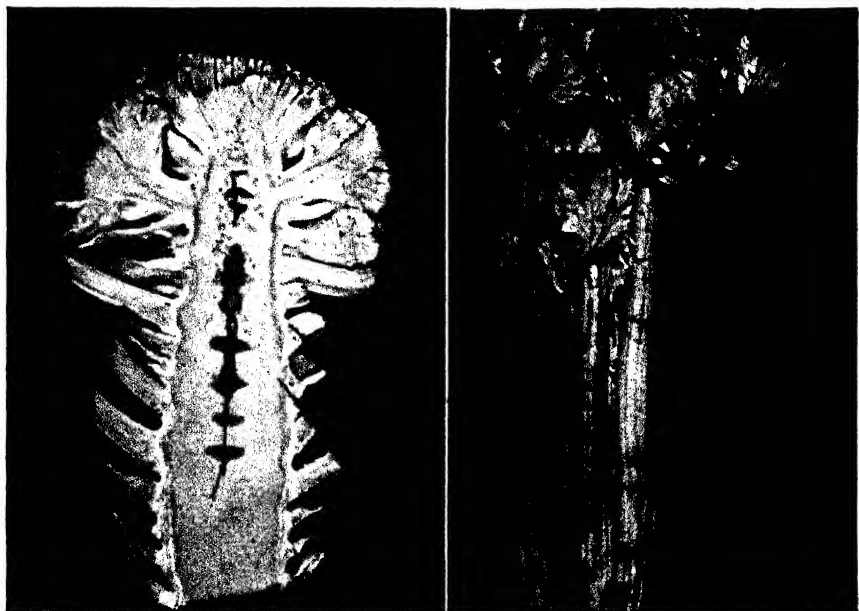


FIG. 69. (Left) In common with many plants cauliflower is very sensitive to boron deficiency. The broken-down tissue in the stalk and brown color of the edible portion of the head are noteworthy. [*Michigan Spec. Bull. 353*, Kodachrome by E. K. Walroth, Eastern States Farmers Exchange.]

FIG. 70. (Right) A deficiency of boron causes "crack stem" in celery. Stalks of this kind have no market value. [*Michigan Spec. Bull. 353*, Kodachrome by E. K. Walroth, Eastern States Farmers Exchange.]

usually be profitable to apply fertilizer long before the symptoms indicating acute starvation actually appear.

*Nitrogen.* The need for more nitrogen is indicated by a light green to yellow appearance of the leaves. As a rule, the older bottom leaves start to turn light green, then get yellow at the tip. The entire leaf may turn yellow, even though the tissues are alive and turgid. In the corn plant the yellowing extends up the midrib of the leaf, with the outer edges remaining green the longest. A nitrogen-starved cucumber may have a small or pointed blossom end; a deficiency of nitrogen may cause the kernels of cereals to become shriveled and light in weight. In fruit trees the early shedding of leaves, death of lateral buds, poor set of fruit, and development of unusually colored fruit are indications of a lack of nitrogen. Extreme nitrogen starvation is most likely to occur in sandy soils or waterlogged soils, although heavy soils with low humus content usually need additional nitrogen.

*Phosphorus.* If phosphorus is deficient, cell division in plants is retarded and growth is stunted. A dark green color associated with a purplish coloration in the seedling stage of growth is a symptom of phosphorus deficiency. Later, plants become yellow. The yellowing is associated with early maturity but is definitely a symptom of phosphorus starvation. Occasionally a pale or yellowish green color develops when the lack of phosphorus inhibits the utilization of nitrogen by the plant. Bronze or purple leaves sometimes are observed at the top of new shoots of phosphorus-starved apple trees. In the absence of sufficient phosphorus, general maturity of the crop and seed formation are usually delayed. With corn, poor pollination frequently is associated with phosphorus starvation. Perhaps the most characteristic symptom of phosphorus deficiency, among plants in general, is the stunted growth.

*Potassium.* A deficiency of potassium usually shows up as a "leaf scorch" in most plants. Corn indicates a need for potassium by a yellowing of the tips and margins of the lower leaves. This coloration does not move up the midrib as with a nitrogen deficiency but gradually spreads upward and inward from the leaf tip and edges. This leaf scorch is frequently mistaken for "burning" or "firing" and is ascribed to a deficiency of moisture during dry weather. When insufficiently supplied with potassium, alfalfa frequently develops a series of white spots near the margin of the older leaves. Sometimes this spotting effect is accompanied by a yellowing of the leaf edges, and at times the leaf margins turn yellow without the formation of white spots. The edges of the leaves finally dry up and curl under. Potato plants indicate a potassium deficiency by a marginal scorch of the lower leaves, and fre-

quently the areas between the veins of potato leaves bulge out, giving a wrinkled appearance. A cucumber, starved for potassium, grows with a small stem end.

**Boron.** Many physiological diseases of plants, such as the internal cork of apples, yellows of alfalfa, top rot of tobacco, cracked stem of celery, and heart rot and internal black spot of beets, are associated with a deficiency of boron. In sugar beets, boron deficiency appears as a stunting and curling or twisting of the petioles, associated with a crinkling of the heart leaves. They have unusually dark green and thicker leaves which wilt more rapidly under drought conditions. The older leaves frequently become chlorotic, and rotting of the beets, starting in the crown, may occur. Girdle or canker of table beets occurs as a cracking of the outer skin of the beets near the soil surface, followed by a breakdown of the root tissue.

**Copper.** There is little evidence to indicate that copper is lacking in soils except those of a high organic content. Growth abnormalities of many plants that are produced on peat and muck soils have been corrected by the application of copper compounds, and in general the copper additions produce better color of such crops as onions, spinach, lettuce, and carrots, increase sugar content of beets and carrots, and improve the flavor of most crops. The need for copper on mineral soils, especially those alkaline in reaction, needs further investigation.

The upper leaves of tobacco plants grown without copper are unable to maintain their vigor, and they wilt badly. The leaves are permanently wilted and do not gain turgidity even in the presence of sufficient moisture. It has also been observed that the amount of seed set is reduced and the seed stalk is unable to stand erect if the supply of copper is inadequate.

**Magnesium.** Magnesium is a constituent of chlorophyll. As with several other nutrient elements, a deficiency of magnesium results in a characteristic discoloration of the leaves. Sometimes a premature defoliation of the plant results from magnesium deficiency. The chlorosis of tobacco, known as "sand drown," is due to a magnesium deficiency. Cotton plants, suffering from a lack of this element, produce purplish red leaves with green veins. Leaves of sorghum and corn become striped; the veins remain green, but the areas between the veins become purple in sorghum and yellow in corn. The lower leaves of the plant are affected first. In legumes the deficiency is shown by chlorotic leaves.

**Manganese.** In the absence of sufficient manganese, tomato, bean, oat, tobacco, and various other plants are dwarfed. Associated with

this dwarfing is a chlorosis of the upper leaves of the plant, and the leaves become spotted. The "gray speck" of oats has been attributed to a shortage of this element in some soils.

It has frequently been observed that the leaves of onions growing in alkaline muck soils become dwarfed and curled during growth and that the bulbs remain immature at harvest time. On similar soils celery becomes yellow; spinach, lettuce, and potatoes are chlorotic and frequently unmarketable.

**Zinc.** Zinc deficiencies frequently have been encountered in Florida. Pecan rosette, the yellows of walnut trees, the mottle leaf of citrus, the little leaf of the stone fruits and grapes, white bud of corn, and the bronzing of the leaves of tung trees are all ascribed to zinc deficiencies. In tobacco plants a zinc deficiency is characterized by a spotting of the lower leaves, and in extreme cases almost total collapse of the leaf tissue may occur.

It should be pointed out that these deficiency symptoms are not always easily diagnosed. Some of them might be mistaken for discoloration or abnormal characteristics produced by diseases, or they may be due to a deficiency of some other element or factor of plant growth. But information concerning these symptoms has been accumulating rapidly and they have become a valuable aid in determining the need for certain nutrients, especially when used in conjunction with tissue and soil tests.

**Pot Tests.** Pot tests with many modifications have been used by various investigators. Essentially these tests consist in filling a number of pots with soil material and adding various fertilizer salts. The need for fertilizer is indicated by the growth of the crop, by amount of dry matter produced, or by analysis of the plant ash. These tests are usually conducted under greenhouse or laboratory conditions and permit the control of moisture supply, temperature, and other factors that cannot be controlled in conducting field tests. Nevertheless, the results have to be interpreted in terms of field conditions if they are to be of value to farmers. This constitutes the chief objection to the method. Regardless of this criticism, pot tests generally are regarded as ranking second to field experiments for determining the fertilizer needs of a soil.

**Field Experiments.** The most reliable method of determining a soil's need for fertilizer is to conduct field tests properly. The reliability of this method lies in the fact that the tests are made under field conditions and on soils in their natural positions. Obviously, the plan of any field experiment is determined by its purpose, but such tests usually are made on a series of plots of equal size by treating the plots in dif-

ferent ways with various kinds and amounts of fertilizer. In the older experiments every third plot is used as a check plot and receives no fertilizer. For most reliable results each treatment should be repeated several times on plots located in different parts of the experimental area. The experiment should be planned so that the data may be treated by statistical methods. Increases in yield of the treated over the untreated plots indicate the need for fertilizer. To carry on long-time field experiments properly requires an enormous amount of time and a considerable expenditure of money. Results obtained from field experiments are invaluable, and such experiments have become the standard method of agricultural experiment stations.

One of the best places to carry on these field experiments is the individual farm. A farmer, for example, can well afford to try fertilizers in an experimental way and to determine for himself if they pay on his particular soils and in his own soil-management system. Each farm is a particular problem in itself.

# 14

## Fertilizers and Fertilizer Materials

Fertilizers, in a broad sense, include all materials that are added to soils to increase the growth, yield, quality, or nutritive value of crops. However, the meaning of the term fertilizer frequently is restricted to apply only to artificially prepared materials containing plant nutrients. Although fertilizers may affect the soil and plant growth in a number of different ways, they are used primarily to increase the supply of available plant nutrients in the soil and also to *balance* the plant-nutrient ratio.

It is customary to speak of fertilizers and fertilizer mixtures as containing nitrogen, phosphoric acid ( $P_2O_5$ ), and potash ( $K_2O$ ) instead of nitrogen (N), phosphorus (P), and potassium (K). The early chemists calculated the results of their fertilizer analyses in terms of  $NH_3$ ,  $P_2O_5$ , and  $K_2O$ , even though these compounds as such were not actually present in the materials. This method, based on the custom of the earlier practice, is used today to express the chemical analyses of fertilizers, with the exception of the values for nitrogen, which have recently been changed and are now expressed in all the states as nitrogen rather than ammonia.

Fertilizers do not consist of the elements nitrogen, phosphorus, and potassium as such, but they are combined with other elements to form either organic or inorganic compounds. Fertilizer materials are classed as nitrogenous, phosphatic, or potassic, depending on whether their principal constituent is nitrogen, phosphorus, or potassium, although some materials can be placed in more than one of these classes. Thus the following subjects necessarily present themselves.

### Objectives

- A. Fertilizer materials supplying nitrogen.
- B. Fertilizers containing phosphorus.
- C. Potassium fertilizers.
- D. Mixed fertilizers.

### FERTILIZING MATERIALS SUPPLYING NITROGEN

The atmosphere is the original source of all nitrogen. In the free state nitrogen is a very inert element. Chemical methods, however, have been perfected whereby this element can readily be combined with others into substances which may be used in the manufacture of fertilizers and for other industrial purposes, and now the possibilities for the production of synthetic<sup>1</sup> nitrogen fertilizer materials are unlimited.

Natural processes have resulted in the accumulation of large quantities of combined nitrogen in soil, coal, nitrate beds, plants, and animals. Relative costs, no doubt, will determine which of the various sources of nitrogen will be drawn on for fertilizer purposes in the future.

Materials used to supply nitrogen are called *nitrogen carriers*, and they may be classified in various ways. A convenient classification is one based on the origin or on the nature of the material. There are thus recognized three general groups of nitrogen carriers: (1) the non-synthetic organic materials, (2) the non-synthetic inorganic materials, and (3) the synthetic nitrogenous materials. Several questions may be kept in mind in studying nitrogenous fertilizers.

#### Questions

1. What are the sources and characteristics of the non-synthetic organic nitrogen carriers?
2. Which non-synthetic inorganic fertilizers are most important?
3. Which of the synthetic nitrogenous fertilizers are most commonly used?
4. Are the various forms of nitrogen equally efficient in increasing crop production?
5. How much of the nitrogen in fertilizers is recovered by crops?

**Non-Synthetic Organic Nitrogen Carriers.** Included in this group are (1) those by-products of animal origin coming from the meat and fish packing and rendering industries, (2) materials of plant origin derived for the most part from vegetable oil industries, and (3) other materials originating from both plants and animals. Several of the more common non-synthetic organic nitrogen carriers with their approximate content of nitrogen, phosphoric acid ( $P_2O_5$ ), and potash ( $K_2O$ ) are given in Table 33. In the early days of the fertilizer industry the carriers of nitrogen were principally the wastes and by-products of other industries, but, since the higher grades of these by-product materials are now used in the manufacture of commercial

<sup>1</sup> Refers to manufactured materials utilizing nitrogen from the air.

TABLE 33

## THE PRINCIPAL NON-SYNTHETIC ORGANIC FERTILIZER MATERIALS

| Material                                     | Nitrogen,<br>per cent | P <sub>2</sub> O <sub>5</sub> ,<br>per cent | K <sub>2</sub> O,<br>per cent |
|--|-----------------------|---|-------------------------------|
| Dried blood                                  | 8.0-14.0              | 0.3- 1.5                                    | 0.5-0.8                       |
| Animal tankage                               | 5.0-10.0              | 3.0-13.0                                    | Small amounts                 |
| Garbage tankage                              | 2.0- 4.0              | 1.0- 3.0                                    | 0.5-1.5                       |
| Process tankages                             | 6.5-10.0              | Variable                                    | Small amounts                 |
| Fish scrap, dried                            | 6.5-10.0              | 4.0- 8.0                                    | Small amounts                 |
| Sewage sludge, ordinary                      | 1.6- 3.3              | 1.0   | Small amounts                 |
| Sewage sludge, activated                     | 4.1- 7.5              | 2.5- 4.0                                    | 0.75                          |
| Cottonseed meal                              | 6.0- 9.0              | 2.0- 3.0                                    | 1.0-2.0                       |
| Bone meals                                   | 0.7- 5.3              | 17.0-30.0                                   | ...                           |
| Castor pomace                                | 4.0- 7.0              | 1.0- 1.5                                    | 1.0-1.5                       |
| Cocoa shell meal                             | 2.5                   | 1.0   | 2.5                           |
| Tobacco stems                                | 1.3- 1.6              | 0.9   | 4.0-9.0                       |
| Sheep or cow manure, dried and<br>pulverized | 1.0- 2.0              | 1.0- 2.0                                    | 2.0-3.0                       |
| Poultry manure, dried and pul-<br>verized    | 5.0- 6.0              | 2.0- 3.0                                    | 1.0-2.0                       |

feeds, cost usually prohibits their use as fertilizers. Organic nitrogen as a rule is more expensive than inorganic nitrogen. In 1913 approximately 52 per cent of the nitrogen in fertilizers was supplied by organic materials, but at present probably less than 5 per cent of the fertilizer nitrogen comes from these sources.

The non-synthetic organic materials have the power to take up considerable water and still retain good drilling qualities. They are useful in preventing hardening or lumping when used in mixed fertilizers, and by virtue of this property they are often referred to as driers or conditioners. The nitrogen in these products is not water soluble, although in most of them it becomes available to crops rather readily under good growing conditions. Biological processes convert the nitrogen into ammonia, which is then changed into nitrates.

**Non-Synthetic Inorganic Materials.** The non-synthetic inorganic materials may be divided into two groups: (1) those obtained from natural salt deposits, such as the Chilean deposits of NaNO<sub>3</sub>, and (2) those secured as a by-product, (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> for example, which is obtained in the process of coking coal. The United States produces more by-product nitrogen than any other country in the world. There are over 85 by-product nitrogen plants (including both coke and gas plants) operating in the United States. This is one of the most important sources of nitrogen for fertilizers.

**Synthetic Nitrogen Materials.** This group includes those fertilizers containing nitrogen taken from the air by any of the chemical nitrogen-fixing processes. The products separate themselves into two groups: (1) organic and (2) inorganic. Urea,  $\text{CO}(\text{NH}_2)_2$ , and calcium cyanamide,  $\text{CaCN}_2$ , belong to the former group, and materials such as  $\text{NaNO}_3$ ,  $(\text{NH}_4)_2\text{SO}_4$ , and  $\text{Ca}(\text{NO}_3)_2$  are placed in the latter group. It should be pointed out that the characteristics of urea and calcium cyanamide are not at all similar to those of the materials listed in Table 33. The nitrogen content of several synthetic nitrogen materials is presented in Table 34.

TABLE 34

THE PRINCIPAL COMMERCIAL FERTILIZER MATERIALS SUPPLYING NITROGEN  
OF SYNTHETIC ORIGIN \*

*Synthetic Inorganic Nitrogen Carriers*

| Nitrogen carrier                      | Nitrogen,<br>per cent | Remarks  |
|---------------------------------------|-----------------------|--|
| Sulfate of ammonia                    | 20.5                  | Also obtained as a by-product from coke ovens                              |
| Nitrate of soda                       | 15.5–16.5             | Also a naturally occurring salt found in Chile                             |
| Calcium nitrate (ANL)                 | 13.0–15.5             | Also called nitrate of lime  |
| Cal-Nitro                             | 16.0–20.5             | Ammonium nitrate mixed with $\text{CaCO}_3$                                |
| Calurea                               | 34.0                  | A compound of calcium nitrate and urea                                     |
| Ammonium sulfate-nitrate              | 26.0                  | A double salt of $(\text{NH}_4)_2\text{SO}_4$ and $\text{NH}_4\text{NO}_3$ |
| Ammonium nitrate                      | 33.0–35.0             | Conditioned to resist absorption of moisture                               |
| Ammonium chloride                     | 26.2                  | Similar in fertilizer properties to $(\text{NH}_4)_2\text{SO}_4$           |
| Anhydrous liquid ammonia              | 82.2                  | Used for ammoniating purposes  |
| Ammonia liquor                        | 24.7                  | Formed when ammonia is absorbed in water                                   |
| Urea-ammonia liquors (UAL—A, B, C, D) | 37.0–45.5             | Solutions of urea in ammonia liquor  |
| Crude nitrogen solution               | 44.0                  | Sodium nitrate and ammonia dissolved in water                              |
| Nitrogen solution II                  | . 37.5                | Ammonium nitrate and ammonia dissolved in water                            |

*Synthetic Organic Nitrogen Carriers*

|                   |           |   |
|-------------------|-----------|---|
| Calcium cyanamide | 21.0–24.0 | Sold under the trade name "Cyanamid"                              |
| Uramon            | 42.0      | This (also Cyanamid N) is classed as non-proteid organic nitrogen |

\* Other nitrogen carriers are listed with the mixed fertilizers in Table 37.

**Relative Merits of the Various Forms of Nitrogen.** Frequent attempts have been made to determine the relative efficiency of nitrogen fertilizers by applying equal quantities of nitrogen per acre, in the various materials, for a given crop. The yield obtained from the use of  $\text{NaNO}_3$  has commonly been used as a standard for comparison. For example, if  $\text{NaNO}_3$  produced 20 bushels of wheat per acre and some other fertilizer produced only 18 bushels per acre, the latter fertilizer would be considered 90 per cent as efficient as the former. Since so many factors, such as temperature and moisture conditions, soil reaction, leaching, kind of crop, and time and method of application, affect the action of any nitrogen fertilizer, relative fertilizing values so obtained may be misleading.

Nitrogen, in the nitrate form, is readily soluble in water and is rather quickly utilized by most crops. Nitrates are easily leached from the soil by rains because of their high solubility and because they are not fixed or held in the soil to any appreciable extent. Although the ammonia form of nitrogen is soluble in water, it is not leached out so readily as nitrates because rather large quantities of ammonia can be adsorbed and held by the soil. A number of different crops can use ammoniacal nitrogen, but most of it is converted to nitrates, owing to the process of nitrification, before plants take it up. In the non-synthetic organic nitrogenous fertilizers the nitrogen exists in the form of complex organic compounds such as proteins, which for the most part are insoluble in water. These substances as such cannot be used by plants but must first be broken down to more simple compounds. This means slower availability of the nitrogen but the furnishing of a more or less continuous supply of available nitrogen throughout the growing season, which is often an important factor in successful plant growth. However, as pointed out elsewhere, the higher cost of nitrogen in these organic materials has greatly reduced the demand for them. The nitrogen in the synthetic organic materials (urea and calcium cyanamide) is largely soluble in water, and under good soil conditions it is rapidly changed to ammonia and nitrates.

Ammonium sulfate and sodium nitrate are among the most widely used nitrogenous fertilizer materials at the present time, and this indicates well the value placed on them. In this group of highly valuable nitrogen materials should also be included many of the newer synthetic products more recently proposed for fertilizer use. It is interesting to note that considerable experimental work has been done on the direct application of anhydrous ammonia; appreciable quantities have been used in this manner in both California and Mississippi. Ammonium

nitrate has also found much favor, but its utilization is limited by lack of production facilities.

**The Recovery of Applied Nitrogen.** The question frequently is asked, how much of the nitrogen applied as fertilizers is recovered by the crop? That is a question which cannot be answered specifically because the amount recovered varies with the kind of fertilizer, soil type, climatic conditions, and plant. But, in general, about 50 per cent of the nitrogen added as  $(\text{NH}_4)_2\text{SO}_4$  or  $\text{NaNO}_3$  will be recovered by the crop. The remainder of the nitrogen is lost from the soil by leaching and erosion, although a small amount may be changed to organic forms by the synthesizing action of microorganisms and retained by the soil. The application of ammonia and nitrate forms of nitrogen in the usual amounts will not result in any great accumulation of soil nitrogen directly, but through their effect in increasing the amount of organic residues the nitrogen of the soil may be increased. Nitrogen of chemical fertilizers cannot be held by soils for any great length of time except as it is fixed in complex organic compounds.

### PHOSPHATIC FERTILIZER MATERIALS

The amount of phosphorus removed from the soils of the United States by erosion, leaching, crops, and grazing greatly exceeds the amount added to the soil in fertilizers, manures, and crop residues. There are no natural processes by which the phosphorus content of our soils can be increased appreciably once it is depleted. In order to increase the phosphorus content of soils, we must buy phosphorus either in the form of fertilizers or in feed. The phosphorus problem, then, differs from the nitrogen problem in that we cannot obtain this element from the air but must eventually buy it if we continue to sell phosphorus from our farms. If purchased in feed for livestock, a portion of the phosphorus contained therein is added to the soil. In general, however, phosphorus must be added mostly in the form of fertilizers which are derived chiefly from phosphate rock. In considering the subject of phosphatic fertilizer materials, the following questions arise.

#### Questions

1. What are the sources of the phosphatic materials, and how are the various carriers prepared?
2. Is the phosphorus of the different carriers equally available to plants?
3. What is meant by the reversion and fixation of phosphates in soils?
4. How can the solubility of phosphorus which is fixed in soils be increased?

**Carriers of Phosphorus and Their Sources.** The principal sources of phosphorus are the natural deposits of phosphorus-bearing rocks, iron ores, and animal bones. The phosphatic fertilizers made from these materials may be classified as (1) the natural phosphates, including materials such as ground rock phosphate and bone meal; (2) the treated natural phosphates, such as bone black, bone ash, superphosphates, calcined phosphates, and calcium metaphosphates; (3) the by-product phosphates, such as basic slag (bone meal could also be placed in this group); and (4) the so-called chemical phosphates, such as ammoniated and nitrated superphosphates, potassium phosphate, potassium metaphosphate, Ammo-Phos, Leunaphos, Nitrophoska, and similar compounds. A list of some of the more common phosphate fertilizers together with their approximate content of phosphoric acid is given in Table 35.

TABLE 35

## THE PRINCIPAL PHOSPHATIC FERTILIZER MATERIALS \*

| Material                                 | Available P <sub>2</sub> O <sub>5</sub> ,<br>per cent | Remarks   |
|--|---|---|
| Rock phosphate                           | 25-35 †   | Effectiveness depends on degree of fineness, soil conditions and crop grown   |
| Superphosphate, ordinary                 | 16-20   | Made by treating ground phosphate rock with H <sub>2</sub> SO <sub>4</sub>  |
| Double, treble, or triple superphosphate | 40-50   | Made by treating ground phosphate rock with liquid H <sub>3</sub> PO <sub>4</sub>   |
| Basic slag                               | 5-20  | By-product obtained in the manufacture of steel   |
| Bone meals                               | 17-30 †   | Includes raw as well as steamed bone meals  |
| Precipitated bone                        | 40  | By-product from manufacture of glue stock from bones  |
| Bone black                               | 32-35   | Bones heated in closed retorts  |
| Bone ash                                 | 32-38   | Bones burned with free access of air  |
| Calcium metaphosphate                    | 60-65   | Prepared by treating phosphate rock with hot, gaseous P <sub>2</sub> O <sub>5</sub>   |
| Colloidal phosphate                      | 18-23 †   | A finely divided, relatively low-grade rock phosphate or phosphatic clay  |
| Defluorinated phosphate                  | 27-33   | Prepared from rock phosphate by heating with water vapor or other defluorinating agent; sold under various names                                |
| Fused or calcined phosphate              | Variable  | Prepared by heating rock phosphate with an alkali or other agent designed to make the phosphorus more available; little fluorine may be removed |

\* See Tables 33 and 37 for other phosphorus carriers.

† Total phosphoric acid rather than available.

*Rock Phosphate.* The United States possesses the largest known deposits of phosphate rock in the world. These deposits are believed to represent 40 per cent of the total known reserves. Extensive deposits of phosphate rock occur in some of the western states (Idaho, Montana, Utah, and Wyoming). Florida also has rather large deposits which are used extensively. Comparatively small deposits in Tennessee are highly developed because of their location. There are small ac-

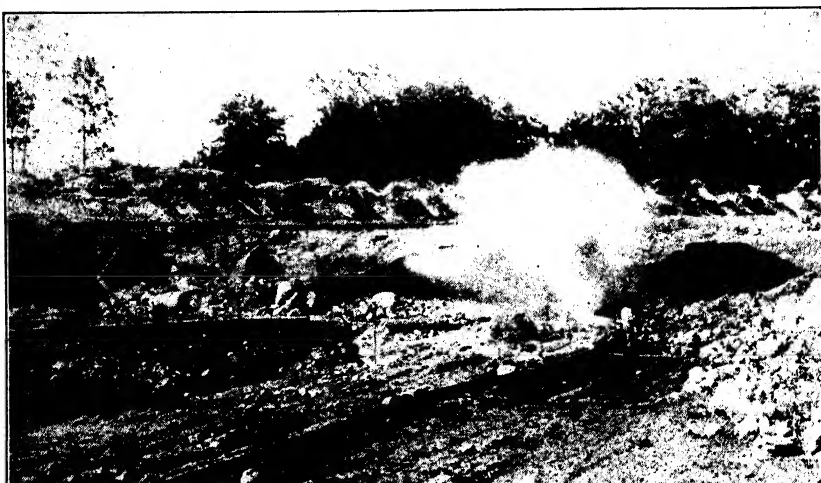


FIG. 71. Rock phosphate is mined from surface deposits by hydraulic pressure. It is then washed and crushed in preparation for acidulation with sulfuric acid. [Courtesy of Agricultural Bureau of American Agricultural Chemical Co.]

cumulations in South Carolina, Kentucky, and Arkansas which are mined little if any.

The principal constituent of American rock phosphate is fluorapatite which may be given the empirical formula  $\text{Ca}_{10}\text{F}_2(\text{PO}_4)_6$ . The formula is sometimes written  $(9\text{CaO} \cdot 3\text{P}_2\text{O}_5 \cdot \text{CaF}_2)$ , which gives the wrong impression because the fluorine is intimately associated with the phosphorus, calcium, and oxygen. This material sometimes is called phosphate of lime and for convenience is frequently expressed as tricalcium phosphate,  $\text{Ca}_3(\text{PO}_4)_2$ . Raw rock phosphate, on account of its insolubility, is not used extensively as a fertilizer. To be effective it is very important that it be finely ground and thoroughly mixed with the soil. Its availability is greatly increased by the presence of decaying organic matter which is believed to exert a dissolving effect on the phosphate

because of the continual production of  $\text{CO}_2$ . Manure or green manure, therefore, aids in making the phosphorus available, and when used under favorable conditions rock phosphate yields reasonably good returns. Rock phosphate should not be used on soils low in organic matter or in those sections of the country where a long and expensive freight haul is necessary.

*Bone Meals.* Various kinds of bone meal are on the market as fertilizers, the principal ones being raw bone meal, steamed bone meal, and precipitated bone meal. Raw bone meal results from the fine grinding of bones without any other treatment. Steamed bone meal is produced by grinding animal bones that have been steamed under pressure. The phosphorus is more quickly available in the latter product because of the finer grinding and the removal of the fat in the steaming process. Precipitated bone is a by-product obtained in the manufacture of glue from bones. Bone meals are expensive forms of phosphoric acid in comparison with superphosphates.

*Superphosphate.* Superphosphate is by far the most important source of available phosphoric acid. More than 90 per cent of all the phosphoric acid used in fertilizers in this country is supplied by superphosphate. It is the most important single fertilizer material, both from the standpoint of total tonnage and from that of quantity of plant food supplied. By treating finely ground raw rock phosphate with approximately an equal weight of sulfuric acid, a superphosphate containing 18 to 19 per cent of phosphoric acid ( $\text{P}_2\text{O}_5$ ) is obtained. This is reduced to the 16 per cent grade by adding filler or is made up to 20 per cent by the addition of treble superphosphate containing 40 to 50 per cent phosphoric acid. (It is possible to produce a 20 per cent product directly by selecting a high-grade raw phosphate rock.) The resulting product of treating raw rock phosphate with sulfuric acid is chiefly a mixture of monocalcium phosphate,  $\text{Ca}(\text{H}_2\text{PO}_4)_2$ , and calcium sulfate,  $\text{CaSO}_4$ , in about equal proportions. The acid treatment of raw rock phosphate thus changes the relatively insoluble fluorapatite into an available form, monocalcium phosphate. The ordinary grades of superphosphate contain, in addition to phosphoric acid, significant quantities of soluble calcium, sulfur, and various impurities, largely silica and clay.

*Treble Superphosphate.* Treble superphosphate (40 to 50 per cent available  $\text{P}_2\text{O}_5$ ) looks very much like the ordinary grades of superphosphate but differs from them in that it contains very little gypsum ( $\text{CaSO}_4$ ). In making these higher-grade superphosphates, liquid phosphoric acid ( $\text{H}_3\text{PO}_4$ ) is used instead of sulfuric acid. In general, the

high-grade superphosphates are just as effective as the ordinary grades and are especially well suited for use in the production of concentrated fertilizer mixtures.

*Fused or Calcined Phosphate.* Rock phosphate has been heated with a large number of materials such as sodium carbonate, gas carrying water vapor, and various acids in order to break up the complex phosphate compound contained in it and thus make the phosphorus more available to plants. Sometimes only a small part of the fluorine is volatilized. In other processes most of the fluorine is driven off. There appears to be a wide variation in the availability of the different phosphates designated fused or calcined.

*Calcium Metaphosphate.* Calcium metaphosphate is a new phosphate fertilizer containing the equivalent of about 62 per cent available phosphoric acid. It is made by treating phosphate rock at a high temperature with gaseous phosphoric acid ( $P_2O_5$ ). The gaseous  $P_2O_5$  is obtained by burning phosphorus (P) which is prepared by heating phosphate rock with silica (pebble) and coke in an electric furnace. The metaphosphate is liquid at the temperatures used and is drawn off as a slag, allowed to cool, and when finely ground is ready for fertilizer use. About one-third of the phosphorus in the metaphosphate comes from the rock and two-thirds from the gaseous  $P_2O_5$ . It has proved to be a satisfactory source of available phosphoric acid for plants.

*Basic Slag.* Basic slag, sometimes referred to as Thomas phosphate, ranks next to superphosphate as the world's leading phosphatic fertilizer. It is produced as a by-product of the iron industry. Phosphorus is contained in certain iron ores, and steel made from them is brittle if most of the phosphorus is not removed. The slag is produced by oxidizing the phosphorus in molten iron by means of a blast of air blown through it. The molten iron is contained in a converter lined with lime, and the oxidized phosphorus combines with this lime. The slag so produced rises to the surface and is drawn off, cooled, and ground so finely that most of it will pass a 100-mesh screen. The phosphorus in slag is soluble in citric acid and is considered available to crops.

*Ammoniated Superphosphate.* This material is made by treating superphosphate with anhydrous or aqueous ammonia. Other materials such as solutions of urea and ammonia, nitrate of soda and ammonia, and ammonium nitrate and ammonia have been used for ammoniating superphosphate. The ammoniation of superphosphate is recognized as an excellent process because the added ammonia may neutralize any excess acidity in the superphosphate. The process makes

use of a cheap source of nitrogen, and the nitrogen content of the fertilizer is increased without appreciably increasing the bulk of material.

*Potassium Metaphosphate, KPO<sub>3</sub>.* If this compound were pure, it would contain 100 per cent K<sub>2</sub>O plus P<sub>2</sub>O<sub>5</sub>. As marketed, however, the product contains 90 to 95 per cent of these two nutrients.

*Other Chemical Phosphates.* In some of the other chemical phosphates such as Ammo-Phos, Leunaphos, Nitrophoska, and diammonium phosphate, the ammonia is combined with phosphoric acid (H<sub>3</sub>PO<sub>4</sub>) instead of with sulfuric acid. These are all high-grade mixed fertilizers, carrying 40 to over 60 units of plant food.

**Relative Availability of the Phosphate Carriers.** The phosphorus compounds contained in phosphatic fertilizers have been arbitrarily placed into three groups, on the basis of their ease of solubility: (1) the water-soluble, (2) the citrate-soluble, and (3) the insoluble. The compounds falling into the first two groups make up the available phosphoric acid content of any particular fertilizer, and the insoluble compounds are considered unavailable. Monocalcium phosphate (the main phosphorus constituent of superphosphate) and the phosphate compounds of potassium and ammonium are soluble in water and are considered readily available. The chemical phosphates are included in this group. Dicalcium phosphate and the phosphorus in defluorinated phosphate, calcium metaphosphate, and precipitated bone are soluble in neutral ammonium citrate, and the phosphorus in basic slag is soluble in citric acid. Therefore the phosphorus in these materials is classed as available. On the other hand, the phosphorus in materials such as bone meal and rock phosphate is neither water-soluble nor citrate-soluble and is thus classified as unavailable, although when these two materials are applied to soils the phosphorus in the former becomes available much more rapidly than the phosphorus in the latter.

The availability of applied phosphorus to crops, however, obviously cannot be determined by the arbitrary solubility tests referred to above. When phosphorus fertilizers are added to soils, the amount that will be or become available for crop use will vary with the type of soil (pH, lime content, organic content), the kind of plant, seasonal conditions, and also with the kind and amount of fertilizer applied. When phosphorus fertilizer is applied to soils, the phosphorus soon combines with various soil constituents to form new compounds, some of which are only slightly available to plants. Therefore, only a portion of the phosphorus added is available for crop use, probably less than 20 per cent in many cases being recovered by the first crop. The processes by which this element is combined with soil constituents are known as phosphate

fixation, and this phenomenon is of considerable practical importance because it affects the efficiency of phosphorus fertilizers on different soil types.

**Reversion and Fixation of Phosphates in Soils.** The changing of phosphorus to a less soluble form is frequently referred to as reversion of phosphate. The changing of monocalcium phosphate to dicalcium phosphate or the changing of either dicalcium or monocalcium phosphate to aluminum or iron phosphate are examples of phosphate reversion. The process of reversion may be considered the same as the fixation of phosphate in soils. Fixation is rapid, and the nature of the compounds formed is obviously determined by the nature of the soil. The phosphorus may be changed to compounds of calcium and magnesium; it may combine with organic matter, be assimilated by micro-organisms, unite with iron and aluminum compounds; or it may be adsorbed by the clay complex in a form such that a part of the phosphorus is exchangeable with other anions.

Soils vary greatly in their capacity to fix phosphate. The organic content of the soil, the pH, the amount of soluble calcium and magnesium, the quantity of iron and aluminum in a reactive form, and the amount and chemical nature of the clay colloids are all important factors affecting the phosphate-fixing power of soils and the nature of the compounds formed. The availability of fixed phosphate depends on the nature of these compounds. For example, when fixed as compounds of calcium and magnesium, phosphorus is generally considered more available than when fixed as compounds of iron and aluminum. The amount applied and the elapsed time since its application are important factors affecting the extent of phosphorus fixation.

The fixation of phosphate in soils is beneficial in that, when fixed, it does not leach out of the soil. But the process is detrimental in that the phosphate may not be readily available to crops, and, furthermore, fixation prevents the penetration of the fertilizer into the soil. As a result, phosphates applied as top-dressings become fixed in the soil surface, and the downward movement into the root zone is very slow. This slow downward movement is an important problem from the standpoint of fertilizing permanent pastures and orchards, for the effects of the phosphate are minimized.

Experiments have indicated that many plants are able to make a more efficient use of phosphorus fertilizer if it is placed in bands to the side of the seed instead of being broadcast. Yields of crops are usually much better from row or hill application than from the use of the same amount of fertilizer applied broadcast because the phosphorus is kept in

a more available condition as it comes in contact with a much smaller amount of soil. The finer a fertilizer is ground, the more rapid and more complete is the fixation of phosphate. Thus the use of granular or pellet forms of fertilizer should result in less fixation and an increased efficiency of the fertilizer.

**Increasing the Solubility of Fixed Phosphorus.** The processes by which the phosphorus in relatively insoluble compounds in the soil becomes available for plant use are not well understood. Because various kinds of organic and inorganic acids, such as  $\text{H}_2\text{CO}_3$ ,  $\text{H}_2\text{SO}_3$ , and  $\text{HNO}_2$ , are produced in the soil, it is assumed that they have a part in making fixed phosphorus more available. The following equation illustrates how such a reaction might proceed; tricalcium phosphate and carbonic acid are used in it.



Phosphate compounds of iron and aluminum frequently are encountered in acid soils, and the application of lime to these soils causes a gradual change of the phosphates to calcium phosphates.

Thus it can be said that the addition of any organic material to the soil which increases the production of carbonic, nitrous, or sulfurous acid will favor the availability of phosphates. Furthermore, the practice of liming acid soils will aid in bringing about the solubility of phosphate compounds, but overliming may temporarily decrease the availability of applied phosphates by causing the soluble phosphates to revert to the tricalcium phosphate form.

### POTASSIUM FERTILIZERS

The supply of potassium in soils is frequently twenty or more times as great as the total supply of either nitrogen or phosphorus. Muck and peat soils offer an exception to this statement, and the supply of potassium in light sandy soils is much less than in heavy soils. In the majority of soils, however, the potash problem is not one of total supply but one of availability, and often the quantity of available potassium is so small that it is necessary to supplement it with potash in the form of commercial fertilizers in order to obtain maximum yields.

The soils in the newer agricultural areas of the United States are, as a rule, not in need of potash, but with the continued use of these soils for the production of crops a deficiency of potash will appear sooner or later. The same situation exists with potassium as with phosphorus in that there are no natural processes whereby this element

is returned in appreciable quantities to the soil. In other words, the farmer must ultimately buy potash because the native supply in the soil cannot furnish crops with sufficient available potassium indefinitely. Before World War I the farmers of the United States were dependent mainly on Europe for their supply of potash, but now all that is consumed as fertilizer in this country can be produced here. In relation to the potash situation, the following questions may be raised.

### Questions

1. What are our important domestic sources of potash?
2. Where are the important foreign sources of potash located?
3. What are the principal carriers of potassium?
4. Are the different carriers of potassium equally efficient?
5. Does fixation of potassium occur in soils?

**Domestic Sources of Potash.** At the beginning of World War I the potash supply from Europe to the United States was cut off and the price advanced over 1,000 per cent, reaching as high as \$500 per ton. This situation did much to stimulate the United States to make every effort to provide an adequate supply of domestic potash. As a result of considerable exploration and research, important sources and deposits of potash have been discovered in the United States, and, as far as this fertilizer element is concerned, this country is self-sufficient.

**Kelp.** This is a seaweed found along the Pacific Coast which contains 11 to 12 per cent of  $K_2O$ . Extraction of the potash is not commercially profitable at the present time.

**Surface Deposits of Rocks and Minerals.** Surface deposits of potash-bearing minerals and rocks are widely distributed over the United States. Some of the more common of these minerals and rocks are (1) alunite (5 to 10 per cent  $K_2O$ ), found in abundance in Utah, Colorado, and Nevada; (2) brown mica (about 10 per cent  $K_2O$ ), found in the leucite-bearing rocks of Wyoming; (3) greensand (6 to 8 per cent  $K_2O$ ), found in New Jersey; and (4) potash shales (8 to 9 per cent  $K_2O$ ), found in Georgia. The potash in all these minerals is insoluble and can be made available only by rather expensive chemical processes. These materials will not be used extensively until more economical methods are developed for the recovery of the potash in available forms.

**Saline Lakes.** These lakes constitute valuable sources of potash in the United States which can be recovered by rather simple chemical processes. Shallow lakes of potassium-bearing brine occur in Nebraska, Utah, Nevada, and southern California. Large quantities of

this element are being extracted each year from the brine of Searles Lake in California. This constitutes one of our most important sources of fertilizer potash. There is also a potash plant at Wendover, Utah.

*Industrial Wastes.* Considerable quantities of potash can be recovered from certain industrial wastes. Sugar beets and sugar cane contain much potassium, and, in the manufacturing of sugar, molasses,



FIG. 72. The porous crystalline mass which fills Searles Lake in California. The brine from this crystal bed is pumped out, and from it are recovered various grades of potash and borax and also lithium chloride. [Courtesy of American Potash Institute.]

and alcohol, a large quantity can be recovered from them. Furnace and flue dusts offer a possible source of potassium. Iron ore and coke, for example, contain different amounts of potash, and many tons of it could be recovered annually from the blast furnaces of the country. The dust from cement kilns also contains considerable quantities of potash.

*Underground Deposits.* In recent years underground deposits of crude potassium salts of commercial importance have been found in the United States, particularly in Texas and New Mexico. The mines located in the vicinity of Carlsbad, New Mexico, supply approximately 85 per cent of our domestic potash. Seventy counties of Texas are underlain at different depths by beds of potassium-bearing salts that vary

greatly in thickness and purity. These deposits are not being exploited at the present time.

*Wood Ashes and Tobacco By-Products.* These products are also potential sources of potash, but, as with other miscellaneous materials, their potash content is too low to justify their extended use. Where these materials can be used locally or where transportation expense

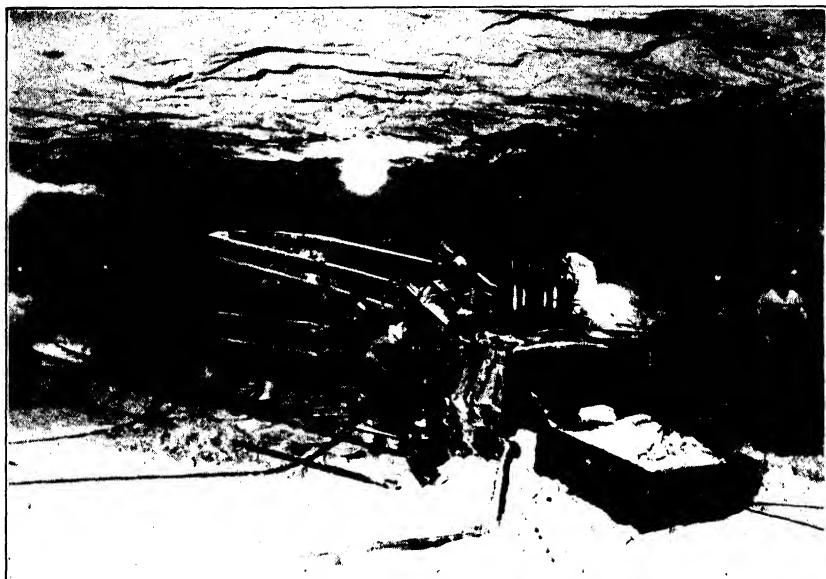


FIG. 73. Crushing and loading crude potash salts in one of the mines at Carlsbad, New Mexico. [Courtesy of American Potash Institute.]

is not excessive, they are of considerable value. At one time (early part of the seventeenth century) wood ashes were exported from the United States for fertilizing purposes.

**Foreign Sources of Potash.** The largest known deposits of materials containing potassium occur in Germany and France. Less extensive deposits are found in Poland, Spain, and Italy. Germany and France have controlled the world's potash market in the past because of their tremendous deposits and because they were the first to develop their potash resources extensively. Furthermore, the deposits are favorably located, being near the large fertilizer-consuming areas of Europe and in a position to take advantage, in peace time, of cheap water-transportation rates to the United States.

The potassium compounds, largely chlorides, in these deep deposits

(approximately 1,200 to 3,500 feet) are more or less impure, being mixed with chlorides and sulfates of calcium, sodium, and magnesium. Most of the salts removed from the mines are ground, dissolved in water, and allowed to recrystallize in order to purify and concentrate them. A small part of the crude products from the mine are finely ground and used directly for fertilizer, or they may be mixed with the refined potash salts.

**Carriers of Potassium.** The principal potash carriers, as used commercially, are potassium chloride (commonly called muriate of potash), potassium sulfate (often referred to as sulfate of potash), potassium nitrate, and manure salts. Manure salt is a semi-refined product, composed chiefly of potassium chloride, magnesium sulfate, and sodium chloride. In some potash-bearing materials, as in wood ashes and molasses waste, the potassium may exist in the carbonate form, but only very small quantities of these substances are used for fertilizing purposes. The principal potassium fertilizer materials and their approximate potash contents are given in Table 36. Potassium chloride is used much more widely than any other forms because of its lower cost.

TABLE 36  
THE PRINCIPAL POTASSIC FERTILIZER MATERIALS \*

| Material                   | K <sub>2</sub> O,<br>per cent | Remarks  |
|----------------------------|-------------------------------|--|
| Potassium chloride         | 48.0–62.0                     | (Muriate of potash)—the most commonly used potassium fertilizer salt                           |
| Potassium sulfate          | 47.0–50.0                     | (Sulfate of potash)—usually sells for a higher price than KCl                                  |
| Potassium carbonate        | 15.0–50.0                     | May be obtained from wood ashes, sugar residue, cottonseed hulls, or other plant materials     |
| Manure salts               | 19.0–32.0                     | Crude potash salts, mostly in the form of KCl  |
| Kainite                    | 14.0–22.0                     | A semi-refined product of potash manufacture, corresponding to the 20 per cent manure salts    |
| Sulfate of potash magnesia | 25.0–27.0                     | A double sulfate of potash and magnesia  |
| Cement dust                | 6.0– 9.0                      | A by-product of cement manufacture   |
| Tobacco stems              | 4.0– 9.0                      | Includes tobacco stalks and tobacco wastes   |
| Hardwood ashes             | 1.5– 8.0                      | The potash exists principally as the carbonate   |
| Cotton-hull ashes          | 10.0–45.0                     | The supply of this material is very limited  |
| Potassium metaphosphate    | 40                            | Prepared by heating KCl with phosphoric acid or P <sub>2</sub> O <sub>5</sub> and water vapor. |

\* See Tables 33 and 37 for other potash carriers.

**Relative Efficiency of the Different Forms of Potash.** All the potash fertilizer salts are soluble in water and are considered readily available. In general, it can be said that there is very little difference in their effects on crop production except in rather special cases. Discrimination is sometimes made against carriers having a high percentage of chlorine for special crops like potatoes and tobacco. The sulfate, or carbonate if available, is usually preferred for tobacco, especially where large amounts are to be added, because a crop of superior burning quality is produced. The muriate is just as efficient as the sulfate for potato production, according to most of the experimental evidence.

**Potash Fixation in Soils.** When a potash salt is added to soil, a part of the potassium is adsorbed or fixed by the soil in more or less unavailable compounds. The amount so fixed is usually directly proportional to the amount of colloidal matter in the soil, being greatest in the heavy soils and least in the light or sandy soils. In general, potassium is not so readily fixed as phosphates but is more readily fixed than most nitrogen salts. The potassium ion, on being adsorbed by the colloids, displaces some other ion such as calcium, magnesium, or sodium. Fixed potassium moves slowly, if at all, in the soil. In light soils the movement may be appreciable, but in heavy soils the movement usually is restricted to the upper few inches of soil. A soil's ability to adsorb and hold potash is of great importance because it serves to decrease leaching and provides for a more continuous supply of available potash. Plants are able to utilize the more easily exchangeable adsorbed potassium.

### MIXED FERTILIZERS

Soils vary greatly in their ability to supply crops with available nutrients, and the mineral requirements of different crops are also quite variable. In order to supply nutrient deficiencies in soils and to meet the various requirements of different crops, fertilizers containing two or more essential elements are prepared in many different grades. They are known as mixed fertilizers and are made by mixing two or more of the separate fertilizer carriers. Thus, some mixed fertilizers contain only two of the three primary plant-nutrient elements and are called incomplete fertilizers. The most commonly used incomplete mixtures contain phosphorus and potassium but no nitrogen. Mixed fertilizers containing all three of the primary fertilizing elements are called complete mixtures. During the year 1948-49, about 69 per cent of the com-

mercial fertilizers sold in the United States were in the form of mixed goods, and most of these mixtures are complete fertilizers. Less than 5 per cent of all mixed fertilizers used are incomplete. In dealing with this subject, the following questions are discussed.

### Questions

1. What advantages are there in using mixed fertilizers rather than using the different materials separately?
2. How are mixed fertilizers prepared?
3. What change has occurred in the strength of fertilizer mixtures used in the United States during the past thirty or forty years?
4. Are low-analysis fertilizers more efficient than either high-analysis or concentrated materials?

**Advantages of Using Mixed Fertilizers.** The use of mixed fertilizers has several advantages over applying the different materials separately. In the first place, the cost of applying a mixture is less than that of applying the various carriers individually; second, mixed fertilizers, as a rule, have better drilling qualities than the individual carriers; and, third, the use of mixed fertilizers requires less care on the part of the farmer to insure a properly balanced fertilizer for his particular conditions than applying the individual materials separately. Mixed fertilizers can be secured from local dealers at such times and in such quantities as the farmer may desire, and it is usually possible to obtain the grade of fertilizer best suited to the particular crops in the locality. Another possible advantage of using mixed fertilizers is the ease with which the acidity of certain fertilizer materials may be controlled. The acidity may be more conveniently and effectively controlled by adding lime to the mixture than by applying it separately in the field.

Mixed fertilizers are made in various grades in order to adjust or bring about a proper balance of the primary fertilizer elements in the soil. If the soil requires only one plant-nutrient element or if the separate carriers can be applied in one operation, obviously the advantages listed above do not hold.

**Preparation of Mixed Fertilizers.** The preparation of mixed fertilizers can be a relatively simple operation, especially if the mixture is to be of low grade, that is, a fertilizer containing a comparatively low percentage of nutrients. It consists essentially in mixing suitable materials in the correct proportion to give the desired grade or analysis. There are many mixing plants in the United States which follow this simple procedure. On the other hand, many of the larger fertilizer factories fol-

low much more involved processes which require careful chemical and temperature control, particularly in the preparation of some of the carriers or materials used in making the final mix.

After the acidulation of rock phosphate in the manufacture of superphosphate, the product is allowed to stand in a large pile for a considerable time to "cure" before being ground. This is also necessary in the preparation of ammoniated superphosphate. If these phosphates are not properly "cured," the chemical reactions involved are not completed and the fertilizer made from them will harden in the bags. Mixtures of superphosphate and potash carriers are often made and stored for a considerable time before being used in the final fertilizer mixture.

There are four types of commercial fertilizer-mixing plants: first, the complete plant that makes its own sulfuric acid, acidulates its phosphate, and then makes the final fertilizer mix; second, the plant that buys its supply of sulfuric acid but grinds its own phosphate rock and acidulates it; third, the plant that makes only its nitrogen carriers and buys its other materials; fourth, the dry-mixing plant, which buys all its materials ready for mixing.

Many of the commonly used grades of fertilizers contain a total of only 1,400 to 1,800 pounds of the different carriers per ton. The other 200 to 600 pounds are composed of some material known as filler. Sand was formerly used extensively for this purpose, but now dolomitic limestone is used to replace a considerable part of the inert material. For example, in 1940 sand and other materials containing no plant nutrients (inert material) made up 12.3 of the 21.0 per cent total filler in mixed fertilizers, whereas in 1945 only 6.8 of the 14.4 per cent filler was classed as inert. Muck and other organic materials are sometimes added to fertilizers in small quantities as filler or conditioners (in 1945, 0.8 per cent). They take up moisture and tend to prevent the mixture from "setting up" or becoming hard on standing.

Fertilizers in storage tend to assume an unfavorable mechanical condition, chiefly the result of "setting up," which is essentially a cementation as in plaster of paris. It may also be due to the surface tension effects of moisture which forms films around the particles of fertilizer.

Superphosphates or other materials which have an excess of sulfuric acid, those which were improperly cured, are greatly improved in their drilling qualities by the use of dolomitic limestone as a filler because it neutralizes any excess acidity.

The drilling qualities of certain mixed fertilizers made from fertilizer salts which take up moisture readily can be appreciably increased by adding an organic filler such as muck.

**Strength of Fertilizer Mixtures.** Fertilizer materials now contain a much higher proportion of the plant-nutrient elements than formerly. When these higher-grade materials are used in mixed fertilizers, it obviously becomes necessary to increase or raise the analysis of the mixture or to dilute it with filler if the analysis is to remain the same. The use of higher-grade fertilizers rather than mixtures diluted with filler affords several economic advantages (see p. 365).

There has been a steady increase in the plant-nutrient content of fertilizers for many years. For example, the percentage of nutrients in fertilizers in the years 1880, 1900, 1930, 1940, and 1949 was 13.5, 13.9, 17.9, 19.8, and 22.55 per cent respectively. This change has resulted because of the demand from fertilizer purchasers for higher-grade fertilizers and the production of more concentrated raw materials.

Various suggestions have been made for grouping or classifying mixed fertilizers according to their plant-nutrient content, but as yet no standard method has been adopted. Differences of opinion exist as to the number of groups that should be made, the limits of each group, and also as to the names that should be applied to each group. The most commonly used classification divides mixed fertilizers into three arbitrary groups, known as low-analysis, high-analysis, and concentrated fertilizers. The first group includes all fertilizer mixtures containing less than 20 per cent of plant nutrients; the second group, those containing 20 to 30 per cent; and the third group, those with 30 per cent or more. It is not to be inferred that the fertilizers belonging to any one of these groups are radically different from those of any other group. This grouping of mixed fertilizers has reference only to the plant-nutrient content and should not be interpreted as indicating in any way the general properties of the particular fertilizers. The composition of several materials containing two or more fertilizer elements is given in Table 37.

**Efficiency of Different Grades of Mixed Fertilizers.** The efficiency of a fertilizer is determined by the uniformity with which it can be distributed and by its quality or its chemical composition. The drillability of a fertilizer is determined by its physical properties, the two most important of which are the relative sizes of particles and their moisture content or hygroscopic properties. A fertilizer is said to be of good quality if it does not cause injury to plants when applied in the usual manner, does not leave any harmful residual effect in the soil, and contains the nutrient elements in proper balance. High-grade mixed fertilizers are now on the market which meet these requirements, and they are being made in granular form to insure good drillability.

TABLE 37

## THE ANALYSIS OF SOME MIXED FERTILIZERS

| Material                   | Available             |   |                               |  |
|----------------------------|-----------------------|---|-------------------------------|--|
|                            | Nitrogen,<br>per cent | P <sub>2</sub> O <sub>5</sub> ,<br>per cent | K <sub>2</sub> O,<br>per cent | Remarks  |
| Potassium nitrate          | 14                    | 0   | 45                            | Occurs only as very small natural deposits                                   |
| Potassium ammonium nitrate | 16                    | 0   | 27                            | Essentially a mixture of KCl and NH <sub>4</sub> NO <sub>3</sub>             |
| Nitrate of soda potash     | 14-15                 | 0   | 10-13                         | A constituent of Chilean salt deposits                                       |
| Potassium metaphosphate    | 0                     | 60  | 40                            | Prepared from KCl and H <sub>3</sub> PO <sub>4</sub> or hot HPO <sub>3</sub> |
| Monoammonium phosphate     | 11                    | 60  | 0                             | Sold largely as a constituent of Ammo-Phos                                   |
| Diammonium phosphate       | 21                    | 53  | 0                             | Stable when made from relatively pure phosphoric acid and ammonia            |
| Ammoniated superphosphate  | 3                     | 16  | 0                             | Analysis will vary with the amount of ammonia added                          |
| Nitrated superphosphate    | 5-6                   | 16-17                                       | 0                             | Superphosphate treated with nitrogen solution II (see Table 34)              |
| Ammo-Phos A                | 11                    | 45-48                                       | 0                             | Largely a mixture of ammonium phosphate and ammonium sulfate                 |
| Ammo-Phos B                | 16.5                  | 20  | 0                             |  |
| Ammo-Phos-Ko *             | 12                    | 24  | 12                            | A mixture of Ammo-Phos and K <sub>2</sub> SO <sub>4</sub>                    |
| Nitrophoska *              | 15                    | 30  | 15                            | There are several different grades   |
| Leunaphos                  | 20                    | 20  | 0                             | Essentially a mixture of diammonium phosphate and ammonium sulfate           |
| Leunaphoska                | 10                    | 10  | 13                            | A mixture of Leunaphos and a potash salt                                     |

\* Trade names for a series of highly concentrated complete fertilizers.

Thus it can be said that a decided increase can be made in the plant-nutrient content of many of the popular low-analysis mixtures without bringing about any marked changes in their chemical or physical properties. In general, a pound of plant nutrients in high-analysis fertilizer mixtures is just as efficient as a pound of plant nutrients in low-analysis fertilizers.

# 15

## Fertilizer Practices

The practice of adding materials to soils to improve their productive capacity dates back several hundred years before Christ. The use of animal manures, marl, chalk, and wood ashes was practiced by the Chinese, Greeks, and Romans more than one thousand years ago. The Egyptians have fertilized their soils with settling from the overflow of the Nile River since the beginning of agriculture in that area. When the white man came to America, he found the Indians putting a fish in each hill of corn. George Washington, in the absence of commercial fertilizers, ran experiments with manure, marl, gypsum, common salt, and green manures.

Although crops have been fertilized in this country for many years, the use of commercial fertilizers dates from about 1830, when Chilean nitrate was first imported. The importation of guanos from Peru, beginning in approximately 1840, did much to acquaint farmers with the general value of fertilizers and to stimulate the rapid development of the fertilizer industry. The production of mixed chemical fertilizers was started in Baltimore in 1850, and the establishment of other plants along the Atlantic Coast soon followed. The use of by-products from meat-packing plants, the residues from plant-oil industries, and scrap from fish canneries and salting plants filled an important place in the early years of the fertilizer industry. Since the development of methods for the fixation of atmospheric nitrogen, the fertilizer industry has changed from a salvage outlet for the waste products of other industries to one of the largest chemical industries.

Farmers in this country were slow at first to use fertilizers because of the large areas of fertile virgin land available for crop production. As the fertility of the older farm soils became depleted, the early farmers moved westward to new territories containing soils of high producing power. But, as time went by and the soils in these newer regions became more depleted, the use of fertilizer increased rapidly.

At the present time the value of commercial fertilizers in farm practice is generally recognized.

Fertilizers should be regarded as sources of plant nutrients for immediate results as well as soil-building materials, although their proper use in adequate quantities may contribute directly or indirectly to an increased supply of nutrients in the soil and an increased humus content. The productive capacity of nearly all soils under cultivation in humid regions can be improved by the use of well-chosen fertilizers, and their use should receive consideration in any good system of soil management. In regard to fertilizer practices the topics listed below are particularly significant and are discussed on the following pages.

### Objectives

- A. Effects of fertilizers on certain soil properties.
- B. Effects of fertilizers on crops.
- C. Laws controlling fertilizer sales.
- D. Home-mixing fertilizers.
- E. The purchase and use of fertilizers.

### EFFECTS OF FERTILIZERS ON SOILS

Soils are complex chemical, physical, and biological systems, and the continued use of fertilizers, especially in large quantities, may produce marked changes in these systems. The beneficial effects of certain fertilizers or the harmful effects of others are no doubt due in part to the influence of the fertilizers on the chemical, physical, or biological properties of the soil. In other words, fertilizers may exert important actions on soils aside from merely increasing the supply of available nutrients. Some of the more important of them are discussed with the idea of gaining a better understanding of the use and function of fertilizers. Furthermore, it is necessary to have a knowledge of some of these effects for the proper interpretation of the results obtained from the application of fertilizers. The questions listed should be answered during the study of this subject.

### Questions

1. What important chemical changes in soils may be produced by the continued use of fertilizers?
2. In what ways may fertilizers affect the physical properties of soils?
3. Are microorganisms in any way influenced by the application of fertilizers?

**Effects on Chemical Properties.** For a number of years the effect of fertilizers on soil acidity has received a great deal of attention. There has been much agitation concerning the manufacture of fertilizers which are neutral, that is, fertilizers which leave neither an acid nor an alkaline residue in the soil. Many fertilizer companies believe the acid-producing effects should be corrected in the fertilizer itself by using ground dolomitic limestone as a filler and by using non-acid-forming materials to supply nitrogen. The amounts of lime so added to soils are too small to be effective in neutralizing an already acid soil, but the lime prevents any possible small increase in soil acidity from the action of the fertilizer. It is much more desirable, therefore, that fertilizer companies use dolomitic limestone as a filler instead of some inert material like sand.

A review of the investigations relative to the effect of fertilizers on soil acidity leads to the following conclusions:

1. The common potassium fertilizers, such as the muriate and sulfate of potash, have no permanent effect on soil acidity.
2. Superphosphates in general will have no permanent effect on soil reaction. Basic slag, bone meal, and rock phosphate have a tendency to correct or neutralize soil acidity.
3. Fertilizers containing nitrogen in the form of ammonia or in other forms subject to nitrification (being changed to nitrate) will produce acidity unless sufficient liming material is present in the fertilizer to neutralize the acid formed. Some experimental fields that have received applications of sulfate of ammonia fairly regularly for many years without being treated with lime have become too sour to grow clover. This effect is more pronounced on the lighter soils such as sands and sandy loams. Fertilizers containing sulfate of ammonia should not be discriminated against, however, since the increased quantity of lime needed to keep the soil in condition for growing legumes in a normal system of soil management is of little practical significance.
4. Nitrogenous fertilizers in which the nitrogen is in the nitrate form and is combined with bases such as sodium or calcium will result, upon being utilized by plants, in decreased soil acidity. Some of the fertilizers in this group are nitrate of soda, calcium nitrate, Cal-Nitro, and Calurea. Calcium cyanamide should be placed in this group although the nitrogen is not in the nitrate form, but the fertilizer carries a rather high content of lime. The acidity or basicity of several nitrogen fertilizers is given in Table 38.

TABLE 38

## EQUIVALENT ACIDITY OR BASICITY OF SEVERAL NITROGEN FERTILIZERS

| Fertilizer Material     | Nitrogen,<br>per cent | Equivalent Acidity or<br>Basicity, pounds CaCO <sub>3</sub> |                              |
|-------------------------|-----------------------|---|------------------------------|
|                         |                       | Per unit of<br>nitrogen                                     | Per 100 lb.<br>of fertilizer |
| Sulfate of ammonia      | 20.5                  | 107   | 110                          |
| Ammo-Phos               | 11.0                  | 107   | 59                           |
| Anhydrous ammonia       | 82.2                  | 36  | 148                          |
| Cal-Nitro               | 20.5                  | 0   | 0                            |
| Calcium nitrate         | 15.0                  | 27 *  | 20 *                         |
| Crude nitrogen solution | 44.0                  | 24  | 53                           |
| Nitrate of soda         | 16.0                  | 36 *  | 29 *                         |
| Urea ammonia liquor     | 45.5                  | 36  | 82                           |
| Calcium cyanamide       | 22.0                  | 57 *  | 63 *                         |
| Cottonseed meal         | 7.0                   | 29  | 10                           |
| Dried blood             | 13.3                  | 35  | 23                           |
| Milorganite             | 6.0                   | 30  | 10                           |
| Garbage tankage         | 2.5                   | 50 *  | 7 *                          |
| Animal tankage          | 6-10                  | 30-30 *   | 15-12 *                      |

\* Basicity.

5. In general, the systematic use of medium to large quantities of high-grade mixed fertilizers at suitable times in the rotation will not appreciably affect soil acidity.

The application of fertilizers may affect, either directly or indirectly, the availability of the soil nutrients. Acid-forming fertilizers in certain soils may decrease the available phosphorus and increase the available potassium. Basic fertilizers may result in an increase in both available phosphorus and available potassium. The addition of basic ions in fertilizers may release various cations from the exchange complex of the soil, making such ions more available and more subject to loss from leaching.

Another chemical effect produced in soils by fertilizers is that of increasing or conserving the reserve supply of nitrogen, phosphorus and potassium. As has been pointed out (p. 350), fertilizers are primarily used to meet the current needs of crops by supplementing and balancing

the soil-nutrient supply, but usually liberal applications of fertilizers ultimately increase the reserve supply in the soil; this is particularly true of phosphorus. An accumulation of either nitrogen or potassium is usually not appreciable, unless the humus content of the soil is increased, because these elements suffer heavy losses by leaching. Any increase obtained in the total reserve supply of nitrogen, phosphorus, and potassium by the fertilizing programs followed by most farmers is insignificant.

**Effect on Physical Properties.** The physical properties of soils are determined largely by the amount and nature of the mineral colloids present and by the content of organic matter. The application of mineral fertilizers does not affect the quantity of soil colloids but may appreciably affect the chemical nature by regulating to some extent the exchangeable cations on the colloids. The kind and amount of exchangeable cations in turn exert a pronounced effect on the physical properties of the clay. For example, the continued use of large quantities of  $\text{NaNO}_3$  may result in producing a soil with poor physical properties. The nitrate radical is adsorbed more by plants than sodium is. The excess sodium combines with carbonic acid ( $\text{H}_2\text{CO}_3$ ) to form sodium carbonate ( $\text{Na}_2\text{CO}_3$ ), which causes soil granules to disperse. In other words, sodium, on being adsorbed by the soil, causes the soil particles to deflocculate or disperse. This may ultimately produce a puddled soil. On the other hand, fertilizers containing a good supply of calcium, magnesium, or potassium tend to promote granulation and thus make for a better physical condition of the soil. In this connection it is to be emphasized that no appreciable accumulation of sodium results from ordinary applications of sodium nitrate ( $\text{NaNO}_3$ ).

Fertilizers may affect the physical properties of soils indirectly through their effect on the quantity of organic matter. In general, fertilizers increase the supply of organic matter by virtue of the fact that they produce more crop residue (roots, stubble, etc.) which is returned to the soil.

**Effect on Biological Activities.** Most soil organisms belong to the plant kingdom, and it is not unreasonable to expect them to respond to applications of fertilizer. All microorganisms require nitrogen, phosphorus, and potassium for their growth and various metabolic functions, and they respond much the same way as do higher plants to applications of commercial fertilizers. Nitrogen-fixing organisms are able to utilize atmospheric nitrogen and consequently are not as a rule appreciably benefited by nitrogen fertilizers, but they do demand an adequate supply of phosphorus and potassium. After the application of

fertilizers, the microorganisms first satisfy their needs, and crops feed on the nutrients left over.

Fertilizers may stimulate biological activity, thereby hastening the decomposition of organic matter, increasing ammonification, nitrification, and nitrogen fixation, and bringing about increases in the solution of soil minerals. Increased acidity as a result of fertilizer applications or the destroying of the granular structure in soil may lead to a decreased biological activity. Certain parasitic soil organisms may be controlled by proper liming and fertilizer practice. It becomes quite evident that many of the observed beneficial effects of fertilizers on higher plants are brought about indirectly through their effect on the soil microorganisms. This feature of fertilizer practice is frequently overlooked.

### EFFECTS OF FERTILIZERS ON CROPS

In addition to their effects on the growth and yield of crops, fertilizers may affect plants in other ways. For example, they may regulate the uptake of certain nutrients by plants, affect the toxicity of certain salts, modify the character of growth, or influence the quality of produce. Several questions that may be raised concerning the effects of fertilizer on plant growth are suggested.

#### Questions

1. Do the ions of fertilizer salts affect the toxicity of other ions or their intake by plants?
2. In what ways may fertilizers affect plant growth?
3. What can be said about the general influence of fertilizers on crop quality and composition?

**Toxicity of Ions and Effects of Fertilizer on Uptake of Nutrients.** A single fertilizer salt may be definitely toxic to plants, but in the presence of one or more other salts there is a tendency for each to counteract the toxicity of the other, thus providing a better nutritive balance. Calcium ions are very effective in decreasing the toxicity of other ions. A fertilizer which is constituted to suit the development of a plant is called a balanced fertilizer, but because of so many variable factors it can be approximated only theoretically. The effect of certain ions on the uptake of certain other ions is a factor which may affect the efficiency of any particular fertilizer. For example, large applications of calcium may depress the uptake of magnesium or potassium by plants if the soil is low in either; high alkalinity may hinder absorption of manganese and iron by plants. It is impossible to forecast all

the reactions that applied fertilizers undergo or the effects produced on other elements, but these various indirect effects must be reckoned with.

**Effect of Fertilizers on Plant Growth.** One of the most important factors affecting the growth of plants is the weather (temperature and quantity and distribution of rainfall), and the response that any particular crop will make to the application of fertilizer is largely governed by the weather, particularly the moisture supply. In seasons when it is necessary to delay planting certain crops because of unfavorable weather conditions, the application of fertilizers may speed up the growth processes of the plant and thus offset somewhat the unfavorable effects of the season. But, looking at the fertilizer-weather relationships from another viewpoint, it is frequently observed that fertilizers in general stimulate early crop growth and, if dry weather prevails about midseason, the fertilizer may result in decreased yields. This is brought about because the soil moisture is more rapidly exhausted by the fertilized crop through increased growth and greater leaf development.

The early growth of a crop should not be taken as a measure of the effect of a fertilizer on yield. At times fertilizers may stimulate early crop growth, but as the season advances this difference disappears, and at harvest no increase is found. Fertilizers may also have little effect on the rate of growth of certain crops, but at harvest a decided increase in yield is noted.

**Effect of Fertilizers on Quality and Composition of Crops.** Some of the more general effects of nitrogen, phosphorus, and potassium on the quality of crops have already been discussed (Chapter 13). The influence of fertilizers on the composition of the mineral matter of plants is exceedingly complex. Their influence is affected by the variety of crop, climate, water supply, and other environmental conditions. The influence which the presence of one element exerts on the absorptive powers of the plant for other mineral elements of the soil or fertilizer also complicates this problem.

Although it is possible to increase to a limited extent the content of mineral constituents of some crops by fertilization, one should not be misled by exaggerated claims for mineralizing human and animal foods. For a more-detailed discussion of this subject see pages 310 and 312.

#### LAWS CONTROLLING FERTILIZER SALES

In the early development of the fertilizer industry in the United States, many low-grade fertilizers were offered for sale in competition with those of higher grade. At that time no fertilizer laws existed for

the protection of the farmer or the purchaser and the honest manufacturer from fraud. There were no effective methods available by which the farmer could be assured of the quality or composition of the different fertilizers in advance of their use. This situation encouraged certain persons to make extravagant claims for materials having little or

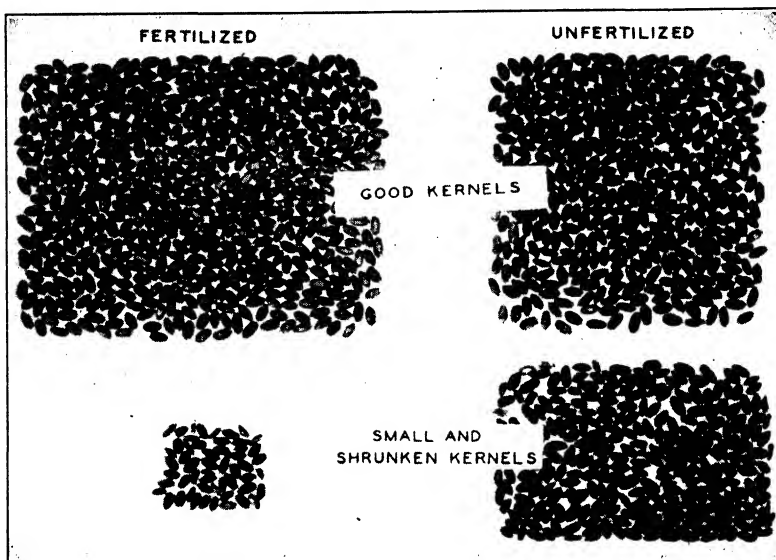


FIG. 74. Properly chosen fertilizer improves the quality of many crops. Left: 20 grams (549 kernels) of wheat grown on fertilized Brookston silt loam soil contained 53 shrunken grains. Right: 20 grams (617 kernels) of wheat from unfertilized soil contained 268 shrunken grains.

no fertilizing value. In order to protect the farmers against fraudulent goods and to protect the reputable manufacturer, nearly all the states have passed laws governing the sale of fertilizers; many states have provided for systems of inspection and analysis. The questions listed below may serve as guides in studying measures governing the sale of fertilizers.

### Questions

1. What is the general nature of the fertilizer laws?
  - a. What order and form are used to express the grade of a fertilizer in most states?
  - b. What information should the guarantee include?
  - c. What is meant by an open-formula guarantee?

2. How are fertilizers inspected, and who makes the analysis?
3. Who is responsible for the enforcement of the fertilizer law?

**General Nature of Fertilizer Laws.** In general, the nature of the laws controlling fertilizer sales in the various states is similar. They all require periodical registration of brands or analyses offered for sale and accurate labeling of the bags or packages. Most of the states require that there be printed on each fertilizer bag, or on an attached tag, the following information:

1. Name, brand, or trademark.
2. Analysis (guarantee) or chemical composition.
3. Net weight of fertilizer.
4. Name and address of manufacturer.

As a rule, fertilizers selling above a certain price require the payment of a license fee for each grade or analysis or payment of a tonnage tax; thus the enforcement of the law is at least self-supporting.

*The Order and Form Used in Expressing the Grade of a Fertilizer.* For every fertilizer material there must be presented a guarantee of its content of nutrient elements although the exact form of stating the guarantee is generally determined by the state in which the fertilizer is offered for sale. In all states the grade is expressed in the order N-P<sub>2</sub>O<sub>5</sub>-K<sub>2</sub>O. The total nitrogen is expressed as elemental nitrogen (N); the phosphorus is expressed as available phosphoric acid (P<sub>2</sub>O<sub>5</sub>); and the potassium is given as water-soluble potash (K<sub>2</sub>O). In all but about ten states the law specifies a minimum plant-nutrient content of mixed fertilizers. As a whole these requirements vary from 14 per cent to 20 per cent, although in one state it is 12 per cent and in one it is 27 per cent.

*Information Needed in the Guarantee.* State laws vary considerably as to the information the guarantee must include. Some do not require all the information that may be desired, while others contain considerable information that is of little value and often is confusing to the farmer. For all practical purposes the analysis should include the following information:

1. Percentage of water-soluble nitrogen.
2. Percentage of water-insoluble nitrogen.
3. Percentage of available insoluble nitrogen.
4. Percentage of available phosphoric acid.
5. Percentage of water-soluble potash.

## Fertilizer Practices

Sometimes the percentages of minor elements may be desired.

The composition of a complete fertilizer, as represented by the analysis, does not give all that is desired in regard to the source of materials. For example, it is possible for a manufacturer to mix high-analysis organic nitrogen compounds with low-grade water-insoluble organics and produce a mixture that may pass inspection in some states. This could be avoided by the use of an open-formula guarantee.

*The Open-Formula Guarantee.* The term *formula* refers to the quantity and grade of stock materials used in making a fertilizer mixture. When both the analysis and the formula are shown on the tag or bag, such a guarantee is called an open-formula guarantee.

**Inspection of Fertilizers.** From time to time during the year the various fertilizers offered for sale are sampled, under official supervision, throughout the state. Fertilizers offered for sale may be sampled by inspectors any time during the year and at any point in the particular state. The samples are sent to the control laboratory and are analyzed to determine whether the goods are up to the guarantee. The results are checked against the guaranteed analysis, and by this means the purchaser is protected from loss through the activities of unreliable companies. The inspection and analysis may be in the hands of the state department of agriculture, of the state agricultural experiment station, or of a state chemist. Once a year, in most states, the findings of the chemists are published in bulletin form, and copies of this bulletin may be obtained upon request by any interested person.

**Enforcement of Fertilizer Laws.** The enforcement of the fertilizer laws in each state is invested in a control official, as indicated above, who is usually associated with either the state department of agriculture or the state agricultural experiment station. When a fertilizer falls below the guarantee (allowing in most cases for a small variation from the guarantee), the manufacturer is subject to prosecution in the state courts. Penalties of varying degrees of severity are levied for failure to come up to the specified guarantee and for other violations of the fertilizer law. The analyses of most commercial fertilizers are generally equal to or greater than the guarantee. Although the farmer is protected by his state laws, the imposition of fines in enforcing these laws is seldom necessary because the publicity derived from the annual publication of the findings of the control agency is usually a more effective means of checking fraudulent guarantees. Not only has publicity been effective in preventing fraud but it is also to the advantage of the fertilizer consumer and of the honest manufacturer to know that the guarantee has received an official sanction. Fertilizers

purchased from outside the state by mail order are not subject to state inspection, and the purchaser is not protected by state law.

### HOME-MIXING FERTILIZERS

Some farmers prefer to buy the raw materials and to mix their own fertilizers, and by so doing they are able to obtain any analysis they may desire. Fertilizers can be mixed satisfactorily on the farm if the farmer is willing to take the necessary time and precautions to do the job correctly. However, the preparation of a fertilizer which will give satisfaction from all standpoints requires considerable technical knowledge. Before undertaking home-mixing one should be familiar with the answers to the questions suggested.

#### Questions

1. What factors should be considered in the home-mixing of fertilizers?
2. What steps are necessary in home-mixing?
3. How are fertilizer formulas calculated?
4. Are there advantages to the farmer in home-mixing?
5. What disadvantages are associated with home-mixing?

**Factors to Consider in Home-Mixing.** First of all, the farmer must decide on the kind and form of nutrients best suited to his soil conditions and crop requirements and then determine the separate materials needed. He must then investigate the cost of home-mixing (including cost of raw materials, hauling, and labor) compared to the cost of purchasing ready-mixed goods in order to determine if home-mixing is profitable. Most of the farmers home-mix because of the possible savings to be had, although occasionally it is practiced in order to secure a particular analysis which cannot be purchased locally.

In deciding on the analysis of fertilizers to be used, the farmer must consider soil conditions such as acidity, texture, organic content, drainage, and soluble-nutrient supply, as well as the cropping system used and the crop to be fertilized. An effort should be made to fit the fertilizer to the soil and crop—no single fertilizer analysis is best for all conditions.

Materials that go into mixed fertilizers cannot be mixed indiscriminately. Certain mixtures may cause the loss of nitrogen, the reversion of soluble phosphates, or they may produce a fertilizer which hardens on standing. It is usually a wise plan not to prepare home-mixed fertilizers until they are needed for immediate use unless great care has

been used in selecting the materials to be mixed and unless satisfactory storage facilities are provided.

Attention should be given to uniformity in mixing the raw materials. Sometimes unsatisfactory results are obtained in the use of a fertilizer owing to a lack of thoroughness in mixing. This would result in an uneven distribution of the nutrients, and under such conditions best results cannot be expected. However, a uniform mixture can be made by the farmer if he puts forth sufficient effort.

**Necessary Steps Involved in Home-Mixing.** The first important step in home-mixing fertilizers is to decide on the analysis. It is then necessary to determine the most desirable kinds of carriers to supply the required amounts of plant nutrients as demanded by the chosen analysis. The formula or recipe (quantity of each carrier required) for the mixture can now be calculated. It is at this stage in the home-mixing process that the farmer should determine if home-mixing will show a profit for that particular year. He will have to compare the cost of the raw materials plus his labor with the price of the ready-mixed goods in the community and then make his decision.

The next step is the actual purchase and mixing of the materials. The mixing process is very simple and requires no equipment other than a shovel, a screen, a tamping or grinding device, a homemade mixer or a tight floor, suitable for mixing, and a pair of scales. Weighing may be unnecessary since the number of pounds of fertilizer contained is shown on each bag, and the bags may be halved or quartered with sufficient accuracy. Any lumps in the fertilizers should be crushed and all material passed through a fine screen before mixing. This insures a more thorough mixing and better drilling qualities.

If the mixture is to be stored for any length of time, it may be necessary to add some sort of drier or conditioner, such as dry sawdust, muck, or soil, in order to prevent lumping. However, in mixing fertilizers on the farm for immediate use, it is usually unnecessary to add a filler, for a proportionately less quantity of fertilizer per acre can be used to supply the amount of plant food desired.

**Calculation of Fertilizer Formulas.** In calculating formulas for homemade mixtures it is necessary to decide first what percentages of nitrogen, available phosphoric acid, and water-soluble potash are desired in the fertilizer mixture and then what materials are to be used in making the mixture.

For example, a farmer wishes to make one ton of a 3-12-12 fertilizer using the following ingredients:

|                   |                                       |
|-------------------|---------------------------------------|
| Ammonium sulfate  | 20 per cent nitrogen                  |
| Superphosphate    | 20 per cent available phosphoric acid |
| Muriate of potash | 60 per cent water-soluble potash      |

The problem is to find out how much of each of these materials is needed. This may be done by use of equation

$$X = \frac{A \cdot B}{C}$$

in which  $X$  equals pounds of carrier required,  $A$  equals pounds of mixed fertilizer required, and, with nitrogen,  $B$  equals the percentage of nitrogen desired in the mixture and  $C$  equals the percentage of nitrogen in the carrier (ammonium sulfate). By substituting these values in the above equation, the result is easily determined.

$$X = \frac{2,000 \times 3}{20}$$

$X = 300$  pounds, the amount of ammonium sulfate required

In a like manner the required amounts of superphosphate and muriate of potash may be determined.

$$X = \frac{2,000 \times 12}{20}$$

$X = 1,200$  pounds, the weight of superphosphate required

$$X = \frac{2,000 \times 12}{60}$$

$X = 400$  pounds, the weight of muriate of potash required

The total amount of materials used in this fertilizer mixture ( $300 + 1,200 + 400$ ) equals 1,900 pounds. It is necessary to add 100 pounds of filler to make a ton of the required mixture.

The quantity of a carrier of a given composition that is required to furnish a given percentage of plant nutrient in a ton of mixed goods, where each nutrient represented in the analysis is obtained from a single carrier, is readily computed from Table 39.

Another type of problem that may need to be solved in home-mixing fertilizers is one in which a part of one or more of the nutrients is derived from two or more materials. For example, using the following

TABLE 39  
TABLE FOR COMPUTING POUNDS OF CARRIER NEEDED TO MAKE 1 TON OF FERTILIZER

| Percentage of Plant Food<br>Desired in Fertilizer | Percentage of Composition of Material (Carrier) |       |       |       |       |       |       |       |       |       |       |       |       |
|---|---|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
|   | 6   | 8     | 10    | 12    | 14    | 16    | 18    | 20    | 22    | 24    | 26    | 28    | 30    |
| 1   | 333   | 250   | 200   | 166   | 143   | 125   | 111   | 100   | 90    | 83    | 41    | 40    | 33    |
| 2   | 666   | 500   | 400   | 333   | 286   | 250   | 222   | 200   | 180   | 166   | 83    | 80    | 66    |
| 3   | 1,000   | 750   | 600   | 500   | 429   | 375   | 333   | 300   | 272   | 250   | 124   | 120   | 100   |
| 4   | 1,333   | 1,000 | 800   | 666   | 571   | 500   | 444   | 400   | 364   | 333   | 166   | 160   | 133   |
| 5   | 1,666   | 1,250 | 1,000 | 833   | 714   | 625   | 555   | 500   | 454   | 417   | 208   | 200   | 166   |
| 6   | 2,000   | 1,500 | 1,200 | 1,000 | 857   | 750   | 666   | 600   | 545   | 500   | 250   | 240   | 200   |
| 7   | 1,750   | 1,400 | 1,166 | 1,000 | 875   | 777   | 700   | 635   | 583   | 292   | 280   | 233   | 226   |
| 8   |   | 2,000 | 1,600 | 1,333 | 1,144 | 1,000 | 888   | 800   | 729   | 666   | 333   | 320   | 266   |
| 9   |   |       | 1,800 | 1,500 | 1,285 | 1,125 | 999   | 900   | 818   | 750   | 375   | 360   | 300   |
| 10  |   |       | 2,000 | 1,666 | 1,428 | 1,250 | 1,111 | 1,000 | 909   | 833   | 416   | 400   | 333   |
| 11  |   |       |       | 1,833 | 1,571 | 1,375 | 1,222 | 1,100 | 1,000 | 916   | 458   | 440   | 366   |
| 12  |   |       |       |       | 2,000 | 1,714 | 1,500 | 1,333 | 1,200 | 1,090 | 1,000 | 500   | 480   |
| 13  |   |       |       |       |       | 1,857 | 1,625 | 1,444 | 1,300 | 1,182 | 1,083 | 542   | 520   |
| 14  |   |       |       |       |       |       | 2,000 | 1,750 | 1,555 | 1,400 | 1,272 | 1,166 | 583   |
| 15  |   |       |       |       |       |       |       | 1,875 | 1,606 | 1,500 | 1,364 | 1,250 | 625   |
| 16  |   |       |       |       |       |       |       |       | 2,000 | 1,777 | 1,600 | 1,454 | 1,333 |

materials, calculate the quantity of ingredients required for a ton of a 2-16-8 fertilizer: one-half of the nitrogen is to be derived from cottonseed meal and one-half from sodium nitrate.

|                   | Analysis |
|-------------------|----------|
| Cottonseed meal   | 6-3-2    |
| Sodium nitrate    | 16-0-0   |
| Superphosphate    | 0-40-0   |
| Muriate of potash | 0-0-60   |

$$X = \frac{2,000 \times 2 \times \frac{1}{2}}{6} = 333.3 \text{ pounds cottonseed meal}$$

$$X = \frac{2,000 \times 2 \times \frac{1}{2}}{16} = 125.0 \text{ pounds sodium nitrate}$$

$$X = \frac{2,000 \times 16 - (333.3 \times 3)}{40} = 775.0 \text{ pounds superphosphate}$$

$$X = \frac{2,000 \times 8 - (333.3 \times 2)}{60} = 255.5 \text{ pounds muriate of potash}$$

333.3 pounds cottonseed meal

125.0 pounds sodium nitrate

775.0 pounds superphosphate

255.5 pounds muriate of potash

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1,488.8 pounds

511.2 pounds filler

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2,000.0 pounds

**Advantages of Home-Mixing.** Usually home-mixing is resorted to because it is economical. Generally, this practice effects some saving, which varies with the analysis to be mixed and the prevailing price of fertilizers, especially if the farmer's labor is not taken into consideration. Home mixtures frequently can be made with farm labor at a saving of 10 to 20 per cent.

Some farmers prefer to mix their own fertilizers so that they can obtain any particular analysis they want; moreover, they can choose their own ingredients. A particular or special analysis may be desired which cannot be purchased locally, and when this unusual condition is present it is advantageous to home-mix.

Perhaps the greatest advantage in home-mixing fertilizer is that it impels the farmer to study fertilizers and their influence on his crops. In other words, home-mixing has a distinct educational value. The farmer must become familiar with the comparative availability and properties of various fertilizer materials; he must spend more time studying his soil and crop needs; and he should understand the residual effects of different fertilizer materials.

**Disadvantages of Home-Mixing.** The practice of home-mixing fertilizers has not become popular in this country largely because farmers do not like to assume the responsibility of buying and mixing the separate materials. Furthermore, it is much easier and more convenient to buy the ready-mixed goods. When it comes time to apply fertilizers, farmers are usually busy and hesitate to take the time required for mixing. If fertilizers are mixed in advance, suitable facilities for storage must be provided.

In comparing the cost of home-mixed fertilizers with that of the factory-mixed goods, the time, trouble, and extra labor involved are frequently overlooked by the farmer. On most farms it is doubtful whether home-mixing should be attempted unless it can be done when other farm work is not pressing. Home-mixing is not economically worth while for most farmers, especially where only a small amount of fertilizer is used, even though it is possible to cooperate with other farmers in purchasing the necessary ingredients (carriers) in larger quantities. Also it is not always easy to obtain the necessary ingredients from local fertilizer dealers in certain areas because these raw materials are not always carried in stock.

The advantages of home-mixing are worthy of serious consideration only by those farmers who use several tons of fertilizer each year. Obviously, fertilizers should be obtained by the method that will give the greatest value for each dollar expended.

#### PURCHASE AND USE OF FERTILIZERS

Good judgment is not exercised frequently by farmers when they buy fertilizers. Too often the price per ton of a fertilizer is the factor governing the purchase rather than the cost per unit of plant nutrients. It is the duty of the purchaser to invest his money in the fertilizer from which he will receive the greatest return per dollar invested in terms of the quantity and quality. However, attention should be called to the fact that, unless the fertilizers are properly used, maximum benefits cannot be expected. It is the purpose of this section, therefore, to

discuss some of the items that determine the price of fertilizers and how they should be purchased and used to obtain the greatest efficiency.

### Questions

1. What charges are included in the retail price of fertilizers?
2. Why should farmers purchase fertilizers on a cash basis?
3. What are some of the more important factors affecting efficiency in the use of fertilizers?

**Price of Fertilizers.** The retail price of a fertilizer is the sum of four charges. First, the price of the plant nutrients contained; second, the transportation charges; third, the profits and selling charges of the dealers, manufacturers, and salesmen; and, fourth, the cost of maintaining and operating the fertilizer factory and the mixing plant and other incidental items. The last three items may be grouped together and called "general expense," which constitutes a more or less fixed charge against each ton. This basic fixed cost is the average cost of manufacturing, bags, transportation, sales, and all other incidentals independent of the plant-nutrient content. The basic cost varies from year to year and also from section to section of the country for any given year. The general-expense item amounts to a large percentage of the total price, particularly of the low-analysis fertilizers, and the basic price would be charged for any mixture handled in the usual way even if it consisted entirely of filler. A source of expense to the user of fertilizer is the purchase of low-analysis mixtures. It can be readily seen that the same amount of plant nutrients will be contained in either 1 ton of 6-18-12 or two tons of 3-9-6, but the consumer would have to pay the general expense charges on two tons if he purchased the 3-9-6 and on only one ton if he chose the 6-18-12. This, of course, would make the plant nutrients in the 3-9-6 mixture much more expensive, even though it cost considerably less per ton. Therefore, it is more economical for a farmer to buy the higher grades of fertilizer, if he makes sure that they contain plant nutrients in proper proportion to meet his needs. One argument advanced in favor of the low-analysis mixtures is the larger quantities of minor or trace elements they may contain. Often, however, it is cheaper to use higher-grade mixtures to which the needed trace elements have been added.

One of the causes of the large overhead on each ton of fertilizer is the making of so large a number of analyses. Every time a mixture of different composition is desired, the mixing machinery must be allowed to run empty and readjustments and arrangements must be made. This procedure results in the loss of much time and in an increase in the ex-

pense. There is no doubt that the number of analyses put on the market in most states could be greatly reduced and that all the needs of farmers and other users of fertilizer would still be supplied.

**Purchasing Fertilizers on a Cash Basis.** Farmers should purchase fertilizers on a cash basis because the cost of fertilizer credit usually is high. Fertilizer credit is different from credit on a piece of farm machinery. If payment is not made on the machinery, the dealer can repossess the implement, but that is not possible with fertilizers. The question of the buyer's good intentions should not be open to discussion, but floods, drought, frost, sickness, or accidents may make payment utterly impossible. These factors combined make the cost of fertilizer credit unusually high. It would be profitable for the farmer to borrow money from a bank or other agency at, say, 6 per cent for a short time in order to avoid payment of the high cost of fertilizer credit.

Most fertilizer companies recognize the advantage of prompt payment and encourage it by rather liberal cash discounts. If a farmer can use carload lots of fertilizer and fertilizer materials and can pay cash, he usually can effect considerable saving by making his purchase direct from the manufacturer, broker, or importer. Cooperative buying of fertilizers by farmers has been employed successfully in many sections, and in this way the small farmer can obtain his fertilizer at as low a price as the large farmer.

**Efficiency in the Use of Fertilizers.** Most soil types, particularly in the humid regions of the United States, respond favorably to the use of well-chosen fertilizers. Fertilizers should not be expected to make up for every shortcoming of the soil and crop, such as poor seed, unadapted varieties of crops, unfavorable weather conditions, poor tillage practices, weeds, poor drainage, bad physical condition of the soil, low organic content, or insufficient lime. All these factors are important, as all affect the efficiency of any fertilizer for any crop grown on any given soil. In other words, the proper use of fertilizer is only one, though a very important one, of the many phases of the scientific management of soil.

The most profitable return from fertilizer is nearly always obtained from those soils that are in the best physical condition for plant growth. The most profitable results from fertilizers cannot be expected when they are used on soils that are too heavy or too light, too compact or too loose, too dry or too wet.

Frequently fields having apparently the same natural characteristics respond differently to the application of a particular fertilizer for the same crop. This condition is often caused by differences in the previous

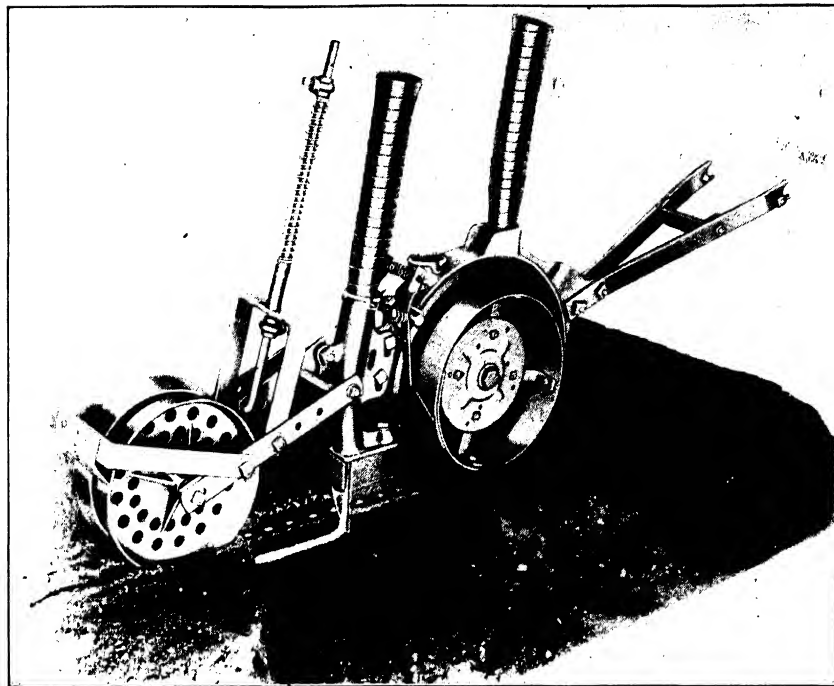


FIG. 75. An excellent device for placing fertilizer at one side of and below the seed. Note also the attachment for varying the depth of planting. [Courtesy of John Deere Co., Moline, Ill.]



FIG. 76. For sugar beets and white pea beans the fertilizer should be placed about 1 inch to the side of and  $1\frac{1}{2}$  inches deeper than the seed. Doubtless the same fertilizer placement will prove advantageous for other row crops.

treatment of the two soils. Variations in past cropping systems, tillage practices, and fertilizer and manure practices bring about differences in the productivity of fields which may be reflected in their response to present fertilization.

Another important factor in the efficiency of fertilizers is the method of applying them; they must be applied in the correct place in relation to the seed. This is probably as important as applying the right kind and amount of fertilizer. No single fertilizer-placement pattern has

been found that is superior for all crops and under all conditions. Almost without exception, however, localized placement of the fertilizer in relation to the seed or plant has been most efficient. This usually means that the fertilizer is placed at the sides of the seed or plant rather than applied broadcast. Obviously, with hay and pasture crops and orchards this method of fertilizer application cannot be followed.

The primary purpose in the use of fertilizers is to produce



FIG. 77. The effect of fertilizer placement on growth of pea seedlings. Left: 300 pounds 4-16-8 in contact with seed. Right: 300 pounds 4-16-8 in bands  $\frac{1}{2}$  inch away from seed and on the same level.

a profit. Space does not permit a detailed discussion of all the factors affecting the efficiency of fertilizers when applied to soils; only a few of the more outstanding factors have been mentioned relative to the judicious use of fertilizers. For more specific information about the best analysis and methods to use, the farmer should refer to fertilizer-recommendations bulletins and consult the county agricultural agent or some other responsible person or agency. Most of the state agricultural experiment stations issue a fertilizer-recommendations bulletin, based on the results of field experiments, which offers the best information available. It may also be pointed out that there is close cooperation between the fertilizer industry and the experiment stations in encouraging the sale of recommended analyses. In several states only the recommended grades are permitted to be sold.

# 16

## Conserving Soil

In the development of a new country little attention is given to conservation. Usually the natural resources which the country affords are present in such quantity that they appear inexhaustible. The limitation of supply is in manpower and in those essentials of life which the country does not produce. Later, when population density has become great, the necessity for conserving resources becomes apparent and frequently acute. Often, as in the United States, the need for conserving soil is one of the last to be recognized. In fact, many people in our country do not yet realize the urgency for conserving soil resources. Our surplus production of a few crops, like cotton, wheat, and corn, tends to obscure the unparalleled waste in soil productivity that is being sustained. Although in a few areas, as in the Southeastern States, Texas, and Oklahoma, some consideration had been given to soil erosion control, the problem as a whole attracted no widespread attention until about 1933. Since that time public interest in the work has developed rapidly.

### Objectives

- A. Soil conservation defined.
- B. Reasons for employing conservation practices.
- C. Controlling erosion caused by water.
- D. Controlling erosion caused by wind.

### SOIL CONSERVATION DEFINED

Before considering in some detail the problem of soil conservation it is advisable to come to an understanding about the interpretation to be given the term; in other words, the natural phenomena, objectives, and practices to be discussed in this chapter. In the early days of the present intensified soil conservation program, great stress was laid on the reduction of soil loss by erosion through the use of mechanical struc-

tures, such as dams of various types and terraces. As the program developed, there came a fuller appreciation of how much may be accomplished by use of cropping systems adapted to the physical characteristics of the soil, and of cropping and planting methods designed to impede water movement and to protect the soil surface from water and wind action. Ultimately the concept of a well-rounded soil-management and farm-management program was evolved. This concept also includes methods of maintaining or increasing the organic content of soil and the supplies of available plant nutrients. In this chapter consideration will be given mainly to the loss of soil through erosion and methods of reducing these losses. The discussion will center around two questions.

### Questions

1. What is the exact, or dictionary, definition of conservation?
2. In what way has expediency modified this definition?

**What the Dictionary Says.** Definitions in standard dictionaries tend to limit the meaning of the word conservation to the idea of preservation. A representative definition reads as follows:

**conser·va·tion** (kōn'sér·vā'shün), *n.* [OF., fr. L. *conservatio*.] 1. A conserving, preserving, guarding, or protecting; a keeping in a safe or entire state; preservation. A step necessary for the *conservation* of Protestantism. *Holloway*. 2. Official care or keeping and supervision, as of a river or forest; conservancy; also, a division, as of a forest, under such conservation. 3. Keeping, as of bees or domestic animals. *Oba*. 4. Making of conserves or preserves, as of fruit.

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Dictionary, Second Edition  
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by G. & C. Merriam Co.

That there has developed an inclination to liberalize the meaning of the word somewhat when applied to natural resources is shown by the following statement from a speech before a conservation congress on September 5, 1910, by William H. Taft: "Conservation as an economic and political term has come to mean the preservation of our natural resources for economical use, so as to secure the greatest good to the greatest number."<sup>1</sup> This tendency to liberalize the interpretation of the word conservation has become quite common in the field of soil science.

<sup>1</sup> Given in *Funk & Wagnalls New Standard Dictionary of the English Language* as a quotation from *New York Times*, Sept. 6, 1910, p. 7, col. 1.

**Conservation as Applied to Soils.** Use of the soil is essential to the maintenance of human life. Accordingly, if we are to apply the idea of conservation to the soil, we must do so with the concept of preservation during utilization. This concept has always prevailed in the philosophy of soil management as advocated by soil scientists and as practiced by large numbers of leading farmers. Members of the Soil Conservation Service, however, have most frequently put this idea into words. One statement which expresses this concept clearly is: "Modern soil conservation is based on sound land use and the treatment of land with those adaptable, practical measures that keep it permanently productive while in use."<sup>2</sup> In this definition there is included the idea of land utilization for the production of food, fiber, and building material. It implies that any given area of land should be studied to determine in what way it can be used for the greatest benefit to mankind, not only for the present but also for the future. The area should then be managed in a way to maintain its usefulness for that purpose.

## REASONS FOR EMPLOYING CONSERVATION PRACTICES

A relatively short trip through virtually any section of our country reveals evidences of soil mismanagement. In the midst of a productive agricultural area many farms will be observed which have been depleted in organic matter content, as evidenced by a lighter color and lower productivity than surrounding soil. In areas of undulating to rolling topography, slopes may be seen which have been denuded of the surface or plow soil, leaving exposed the lighter colored subsoil. And, again, many instances are seen of sandy soils which have been cleared of their forest cover, cropped for a few years until the virgin fertility was exhausted, and then abandoned to become covered with weeds and brush or to be blown about by the wind.

Another evidence of the need for soil conservation is the fact that in most areas of our country yields of commonly grown crops have not increased in proportion to our knowledge of insect and disease control methods, improvements in crop varieties and seed quality, and the use of lime and fertilizer. Decrease in soil productivity is responsible, at least in part, for the failure of crop yields to keep pace with agronomic knowledge.

As pointed out in Chapter 20, many nations have inadequate food supplies, and a greatly increased food production will be needed to

<sup>2</sup> H. H. Bennett in *Soil Conservation* for September, 1949.

feed the expected world population in 1960. This situation affords another reason for giving careful attention to soil preservation. Answers to the following questions should be found in the discussion.

### Questions

1. What is the difference between geological and soil erosion?
2. Is erosion a natural process?
3. Have soils in the United States suffered seriously from erosion?
4. Is the supply of good soil in the United States limited?
5. How much productive soil is needed?
6. Does erosion cause damage in addition to loss of surface soil?
7. Are streams and harbors obstructed by erosion?
8. Is the capacity of reservoirs decreased?

**Geological vs. Soil Erosion.** Erosion which takes place under natural conditions, that is, when the land surface and native vegetative cover have not been disturbed by man's activities, is called natural or *geological erosion*. On the other hand, when timberland is cleared or grassland is broken up, processes of erosion are accelerated and we have unnatural or soil erosion. It is sometimes said that, whenever erosion is speeded up as a result of man's activities so that it removes all or part of the topsoil, we call the process *soil erosion*.

Geological erosion is a relatively slow process under many conditions, and soil formation may keep pace with the removal of the surface soil. Soil erosion, on the contrary, is very rapid when environmental factors favor erosion. For example, at the Statesville, North Carolina, Experiment Station, land planted to cotton year after year suffered an annual loss of 31.22 tons of soil per acre. The same kind of land, however, under natural conditions lost only 0.002 ton of soil per acre annually by geological erosion. Nevertheless, over long periods of time geological erosion moves very large quantities of soil material to lower levels, as pointed out in the following section.

**Geological Erosion a Natural Process.** Geological erosion is a natural process which tends to bring the earth's surface to a uniform level. Whenever one part of the surface of the globe is elevated above the surrounding portions, erosion immediately begins the work of leveling off the high land. The leveling process may result in a very rough topography in the early stages by the cutting of gullies or of canyons in a mountainous region, but the ultimate result is a comparatively level surface. Evidence of this geological process is seen in peneplains, mesas, valley fills, alluvial plains and deltas, extensive deposits of wind-laid material, and numerous other geological formations. Some concept

of the extent of erosional activity may be gained from the fact that the Appalachian Mountains are said by geologists to be only about one-half of their original height. There are also remnants of other mountain ranges in the United States which are now merely rough land or low hills.

**Damage Done to Agricultural Land by Soil Erosion.** The damage done to farm lands in the United States since they were occupied by the white man can only be estimated roughly. However, from data supplied through soil surveys, erosion surveys, and other inventory methods it has been suggested that erosion has ruined or seriously damaged some 282,000,000 acres in the United States. A far larger area, possibly three-quarters of a billion acres, has suffered the loss of a greater or lesser portion of its topsoil.

Of land used for crop production, it is estimated that around 50,000,000 acres have been rendered useless and a like amount is approaching that condition. An additional 100,000,000 acres, although still being cultivated, have lost one-half or more of the surface soil, and on a similar acreage erosion is carrying on its insidious work of destruction. Dr. H. H. Bennett, chief of the Soil Conservation Service, has expressed the opinion that erosion control "is the first and most essential step in the direction of correct land utilization on about 75 per cent of the present and potential cultivated area of the nation."<sup>3</sup> It may be truthfully said that there are very few farms on which erosion by water or wind is not causing some damage to soil or crops.

In considering the damage done by erosion one should keep in mind the fact that a large share of the soil lost by this process is the surface or plow soil. It is this soil layer which contains the highest percentage of many of the essential plant-food elements in an available condition. Plant scientists consider that most crops obtain the major portion of their mineral nutrients and nitrogen from this part of the soil. Furthermore, studies of the soil eroded from fields in many parts of the country have shown these to be made up largely of the finer soil particles (clay, silt, very fine sand, and humus). These particles contain a higher percentage of several of the plant nutrients than do the coarser particles.

**Acreage of First-Class Land in the United States Limited.** In colonial days and for many years afterward, the supply of productive soil in the United States was considered unlimited. With the spread of people to all corners of our country and the rapid increase in pop-

<sup>3</sup> By permission from *Soil Conservation*, by H. H. Bennett, copyright, 1939, McGraw-Hill Book Co., Inc.

ulation there came to soil scientists and agricultural economists the question, how much land suitable for crop production is there in the United States? Coupled with this query was the desire to know how the land should be classified from the point of view of productivity. In other words, how much of our land is really first-class, and how much is mediocre and poor in quality? An early attempt to classify land on this basis was made by the former Bureau of Chemistry and Soils of the United States Department of Agriculture. Results were published in the Report of National Resources Board, 1934. According to this compilation the land area of continental United States was classified as 5 per cent excellent, 11.1 per cent good, 18.1 per cent fair, 19.1 per cent poor, and 46.4 per cent unfit for production of general crops without irrigation, drainage, or fertilization. These figures show an alarmingly small amount of good and excellent land.

Further studies along this line have been made under the leadership of Dr. C. E. Kellogg, chief of the Division of Soil Survey in the Bureau of Plant Industry, Soils, and Agricultural Engineering, and also by the Soil Conservation Service. The results are discussed in Chapter 20.

It is evident from these results that the supply of good farm land in the United States is definitely limited, and, if we are to feed and clothe adequately the population expected in the years to come, definite steps must be taken to conserve our farm soils.

**Changes in Acreage Requirements per Capita.** It is difficult to calculate the acres which are required to produce the food needed by each citizen of our country. The productivity of the soil varies with natural fertility and management practices. Advances in agronomic knowledge dealing with crop varieties, insect and plant disease control, and cultural practices contribute to production. The nature of the diet is also a factor. Fewer acres are needed when the diet is made up largely of vegetables and grain crops than when a considerable quantity of meat is consumed. As a general average, perhaps it is safe to say that from 1.5 to 2.0 acres per person are needed for food production exclusive of grazing land. It is evident, therefore, that as the population increases the demand for land increases quite rapidly.

**Erosion Damage Not Confined to Soil Loss.** Erosion by water opens the way for at least five types of loss or damage.

1. The loss of the water causing the erosion. It might have been useful in crop production had it entered the soil instead of running off over the surface.

2. The soil carried away by erosion frequently ceases to be of value in crop production, and, furthermore, the remaining soil, denuded of the surface or plow soil, is much decreased in productivity.

3. The soil carried away frequently causes much damage. Especially during gully formation, a layer of infertile subsoil may be deposited over an area of productive soil, thus greatly reducing the crop-producing power.

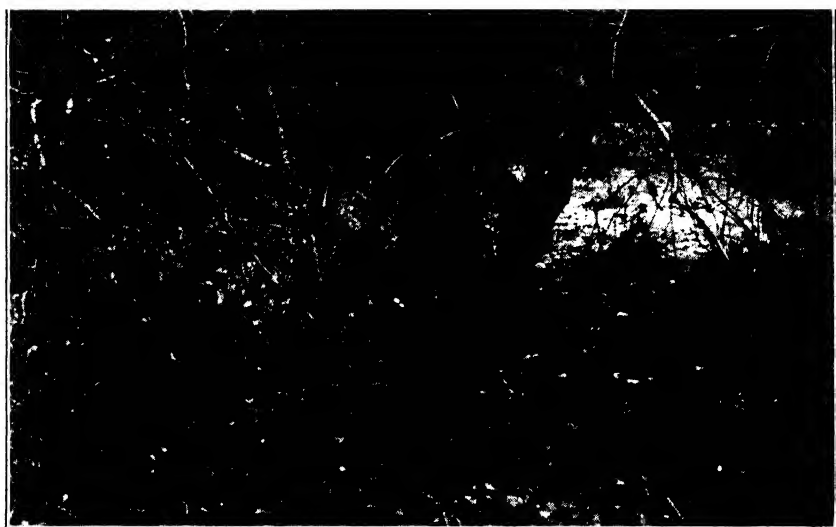


FIG. 78. Soil eroded from poor land may greatly damage crops growing on good soil. This cherry tree is being buried by coarse sand washed from adjacent slopes. [Courtesy of Soil Conservation Service.]

4. Another damage resulting from gully formation is the cutting up of fields into irregular pieces. As these gullies get too deep to cross with farm implements, great inconvenience and loss of efficiency in cultivating the land and planting and harvesting crops result.

5. The soil removed through erosion may be deposited in streams, harbors, and reservoirs, thus increasing floods, impeding navigation, and reducing water-storage capacity.

**Need for Cleaning Streams and Harbors.** Sediment in ditches and streams obstructs the flow of water and so impedes drainage of farm land, increases the hazard of floods, and reduces transportation facilities in navigable streams. The cleaning of ditches, streams, and

harbors involves a large expenditure in our country much of which must be paid for by taxation. An example is the experience with the Saint Joseph river. This stream rises in southern Michigan, loops down into northern Indiana, and then turns north, entering Lake Michigan between Benton Harbor and Saint Joseph. Its course lies in well-developed farm land. According to the 1948 report of the Chief of Engineers, U. S. Army, some 84,563 cubic yards of sediment were removed from the harbor at a cost for dredging of \$42,531.01. This sediment consisted of material washed from Michigan and Indiana farms. It should be pointed out also that an additional large tonnage of the fine soil particles, which also came from the farms in the drainage basin of the river, was carried into the lake in suspension.

The amount of sediment carried by various streams is enormous, as is shown by the estimates made by the U. S. Geological Survey (Table 40).

TABLE 40  
TONS OF SEDIMENT CARRIED BY SEVERAL RIVERS

| River       | Tons per Year | River     | Tons per Year |
|-------------|---------------|-----------|---------------|
| Hudson      | 240,000       | Savannah  | 1,000,000     |
| Susquehanna | 240,000       | Tennessee | 11,000,000    |
| Roanoke     | 3,000,000     | Missouri  | 176,000,000   |
| Alabama     | 3,039,000     |           |               |

**Rapid Sedimentation of Reservoirs.** The period of service of reservoirs, formed by damming streams, is greatly reduced by sediment carried into the stream by erosion processes. Such reservoirs are often essential to large numbers of people either as sources of electric power or water for irrigation or household use. The reduction of their storage capacity through sedimentation accordingly becomes a matter of much concern.

During the period 1934-38 the Soil Conservation Service made a survey of a large number of reservoirs in the United States. Representatives of the U. S. Bureau of Reclamation assisted in some of the surveys to determine the amount of sedimentation which had taken place. A summary of the data for 24 reservoirs in southeastern United States with an average age of some 15.3 years showed that they had lost an average of 20.8 per cent of their storage capacity. An extreme example of the loss of storage capacity as a result of erosion is furnished by the Washington Mills reservoir at Fries, Virginia. In the course of 33.5 years 82.69 per cent of the storage capacity had been lost.

A study of Lake Decatur, municipal-water-supply reservoir of De-

catur, Illinois, showed that it had lost 1.0 per cent of its storage capacity annually between the date of construction (1922) and 1936. Between 1936 and 1946 the annual rate of sedimentation had increased to 1.2 per cent of the capacity. Also, in a land-utilization project area near Pierre, South Dakota, the annual rate of silting of stock ponds was found to vary between 1.10 and 5.56 per cent.

The Roosevelt Dam, which supplies water for electric power and for irrigation of the great Salt River Valley in Arizona, lost some 6.9 per cent of its storage capacity through sedimentation during the first 24 years of its existence. Likewise, the Elephant Butte Dam on the Rio Grande River in New Mexico decreased in storage capacity about 16.59 per cent during the period between 1915 and 1947. Such losses of storage capacity are serious matters, especially when the supply of water is scarcely adequate to meet demands.

#### CONTROLLING EROSION CAUSED BY WATER

It is difficult if not impossible to differentiate the erosion damage caused by water from that brought about by wind. In areas of high rainfall there are periods during which soils of sandy texture blow badly when not protected by vegetation. Also, the quantity of material moved by wind from the surface of heavy-textured soils is surprising. On the other hand, in regions of low precipitation there is much erosion by infrequent but torrential rains, although wind is the predominant agency of erosion. Because populations are more dense and interest in agriculture more general in regions of moderate to heavy rainfall, erosion caused by water has attracted the attention of more people than has wind erosion. As a result, more effort has been expended in devising means of controlling water erosion.

In discussing erosion control it is well to remember that this phenomenon really involves *two* processes: (1) the loosening or detachment of soil particles or groups of particles from the main body of the soil, and (2) the transportation or removal of these particles or particle groups from their position. These ideas are contained in the definition of soil erosion taken from *Soil Conservation* for February, 1946: "Soil erosion is a process of detachment and transportation of soil material by erosive agents." Three fundamental principles which guide many of the methods employed to reduce erosion by water are: (1) protect the soil surface from impact of raindrops; (2) prevent water from concentrating and moving down a slope in a narrow path; and (3) impede the movement of water so that it moves down a slope slowly. Several

questions may be helpful in studying methods of controlling erosion caused by water.

### Questions

1. What types of water erosion are there?
2. Can gully formation be controlled?
3. How does rain influence the physical condition of the soil?
4. Does type of storm influence erosion?
5. To what extent does slope affect erosion?
6. Can soils be made to absorb more water?
7. Can erosion be reduced by crop rotation?
8. Should land characteristics be considered in planning a cropping system?
9. What is meant by strip cropping?
10. Is terracing a practical means of erosion control?
11. Under what conditions are sod waterways useful?

**Types of Water Erosion.** Erosion by water may be divided into four categories, namely, splash, sheet, rill, and gully. Strictly speaking, sheet erosion refers to the quite uniform removal of soil from the surface of an area in thin layers. For sheet erosion alone to occur it is necessary that there be a virtually smooth soil surface, which is seldom the case. Usually a soil surface designated smooth contains small depressions in which water will accumulate. Overflowing from these at the lowest point the water cuts a tiny channel as it moves down the slope. Duplicated at innumerable points, this process presently creates a surface cut by a multitude of very shallow trenches which are called rills. None of these may grow to appreciable size or depth, and so the surface soil is rather uniformly removed from the field. Accordingly, sheet erosion and rill erosion work hand in hand and the combined process is usually called sheet erosion, as distinguished from gully formation.

Most fields are sufficiently irregular to cause water to move into natural depressions as it flows down the slope. This action causes the erosive force of the water to be concentrated on a narrow surface; thus the topsoil is quickly removed and the subsoil begins to wash away. In a relatively short time a trench of appreciable size is cut, and gully formation is under way. Although some gullies never become very large because of the lack of height or length of the slope, others have grown to great depth and width.

A type of erosion which received little attention until recent years is the splashing or scattering of the smaller soil particles by the impact of raindrops. At first thought this action seems trivial, but, when con-

sideration is given to the large number of raindrops that strike a square inch of soil surface during a one-hour rain and the force with which they strike, it is seen that the net effect in loosening and moving soil particles may be considerable.

*Sheet Erosion Insidious.* Rills are obliterated by the first cultivation of a field even by a harrow, and, in consequence, attention is not attracted to sheet erosion. A farmer may be conscious that the surface soil is unusually deep at the foot of a slope, but it does not occur to him to worry about it. So sheet erosion goes on largely unnoticed until the yellow or brown subsoil begins to appear on the steepest part of the slope and crop growth at that point is thin and unthrifty. Even then many farmers take the occurrence as a matter of course and do not feel the need of putting into practice erosion control methods. A little extra manure is spread on that part of the field, and farming goes on as usual. The very fact that sheet erosion fails to attract attention accounts for the vast damage done by this process. Also, the occurrence of sheet erosion on slopes which are quite gentle leads one to overlook the loss of soil. Although far less noticeable, the total damage done by sheet erosion greatly exceeds that caused by gullying. On many farms sheet erosion is likely to occur on the best land.

*Gully Erosion Spectacular.* Although sheet erosion may pass unnoticed by the average observer, gullies attract immediate attention. They disfigure the landscape and give the impression of land neglect and soil destruction. Not only do gullies result in soil loss but also, as previously mentioned, the eroded material is usually deposited over more fertile soil at the foot of the slope. Also, fields dissected by gullies offer many problems in farming operations.

Gullying proceeds by three processes: (1) waterfall erosion, (2) channel erosion, and (3) erosion caused by alternate freezing and thawing. Usually more than one process is active in a gully. Water falling over a soil bank undermines the edges of the bank, which then caves in, and the waterfall moves upstream. This process produces U-shaped gullies, particularly if the underlying soil material is soft and easily cut. Gullies which are V-shaped are produced by *channel erosion* through the cutting away of the soil by water concentrated in a drainageway. This type of gully usually forms when the underlying soil horizons are of finer texture and more resistant to erosion than the surface horizons. Soil loosened from sides of gullies by alternate freezing and thawing sloughs off and is then carried away by heavy rains. Gullies are sometimes classified as small if they are less than 3 feet deep, medium-sized if from 3 to 15 feet deep, and large if over 15 feet in depth.



FIG. 79. This Michigan gully was formed in 25 years by an improper terrace emptying out into a road. [Courtesy of Soil Conservation Service.]

**Concentration of Water to Be Avoided.** As pointed out, gully formation results from water concentrating in a narrow channel to flow down a slope over unprotected soil. Careless and often seemingly trivial mismanagement practices in the handling of land are most frequently responsible for the starting of gullies. It is much simpler and less expensive to prevent the concentration of water before it flows down a slope than it is to stop gully formation when the process is well established. Some of the procedures which have been found effective in preventing water from accumulating in a channel are based on two principles: (1) cause the adsorption of excess water in the soil of the area which feeds the channel, and (2) divert runoff water so that it runs around rather than into the drainageway. Both in curing gullies and in preventing them, attention should be given first to absorption of the excess water. When these measures are not sufficient to prevent water concentration, use should be made of the second principle.

Some of the practices used in increasing absorption of rainfall are: (1) a good crop rotation, including ample use of sod crops and cover crops, (2) permanent pasture when necessary, (3) strip cropping, (4) contour cultivation alone or in combination with terracing. Varying

types of contour furrowing and listing make long trenches which hold large quantities of runoff water until it can be absorbed. Terraces as well as diversion ditches carry excess water around existing or prospective gully heads. The safe disposal of such diverted water is a most important problem.

Sometimes it is necessary to convey part of the runoff through a gully. This can be done safely if vegetation can be established in the gullies or if suitable mechanical structures are built at critical points to supplement the vegetative cover. Some small gullies may be stabilized by plowing in the sides, sodding the heads, and placing sod strips across them at suitable intervals. Many practices which cannot be mentioned in this brief discussion are in use to fill or stabilize gullies effectively. The fundamental principle behind most of them, however, is the establishment of a vegetative cover of suitable type.

**Damage Due to Impact of Raindrops.** Soil particles must be dislodged from their resting place before they can be moved. This is one part of the erosion process. Except in very sandy soils, dislodgment usually involves the breaking down of small clods or soil granules. The impact of raindrops on unprotected soil is one of the main forces of particle dislodgment. The kinetic energy of a raindrop is proportional to the product of its mass and the square of its velocity. There are data which indicate that the velocity of falling rain is of the magnitude of 20 miles per hour, and that a 2-inch rain on an acre would have 6,000,000 foot-pounds of kinetic energy. This is sufficient energy to raise a 7-inch layer of soil 3 feet. There are, of course, great variations in the drop impacts of different storms. Studies by W. D. Ellison, of the Soil Conservation Service, show that these variations may result in a difference in quantity of soil moved per acre of from less than 1 ton to more than 100 tons. Differences in the characteristics of different soils also cause vast variations in the quantity of soil splash. Measurements have shown that soil particles may be splashed to a height of over 2 feet and may be moved more than 5 feet horizontally on level surfaces. Soil particles thrown into the air will naturally fall downhill on steep slopes, and hence splash from raindrops may result in considerable soil movement. Probably, however, the dislodgment of soil particles so that they are carried by water moving on the soil surface is a more detrimental effect of falling raindrops. Another type of soil damage caused by the impact of raindrops is discussed next.

**Disintegration of Soil Granules by Raindrops.** Rain soaks rapidly into soil composed mainly of large granules or of small clods. As long as these structural units remain intact, the spaces between them



(a)



(b)



(c)

FIG. 80. How raindrops cause soil erosion by "splash." (a) A drop just before it strikes the ground. (b) Just after the drop struck the soil surface. (c) What happens when the drop "explodes" like a small bomb scattering fragments of soil in all directions. [Courtesy of Soil Conservation Service.]

are sufficiently large to offer little resistance to the entrance of water. However, water soon softens the binding materials which hold the particles together, and then the granules and clods disintegrate. The impacts of the raindrops hasten this disintegration and also bring about a separation of the finer particles by causing them to splash into the air. This process results in an accumulation of the very fine particles on the soil surface. Evidence of this phenomenon is seen in the thin layer of fine sediment which tends to crack and curl up on the soil surface as it dries after a heavy rain. These minute particles fill any spaces between larger particles and result in a "skin" or layer over the soil which permits only very slow entrance of water. In consequence, a high percentage of the rainfall runs off over the soil surface during hard beating rains on sloping land. As would be expected, this "sealing" of the soil surface is more pronounced on soils containing a high percentage of colloidal clay but also occurs to a surprising extent on many sandy loams.

If the granules have a high stability, they do not disintegrate so readily and hence the soil may not seal over so badly in storms of moderate intensity. Soil granules produced by growing sod crops seem to have a greater stability than those produced by freezing and thawing, or by wetting and drying. Likewise, a high organic content in soils appears to keep them in a better condition for the intake of moisture. Needless to say, any soil covering such as crop residues and mulches, which protects the soil from the impact of raindrops, will maintain it in the very best water-receptive condition.

**Torrential Storms Most Detrimental.** Strange as it may seem, water erosion frequently causes more damage in regions of low rainfall than in parts of the country having a moderate annual precipitation. This results from the fact that, although rain is less frequent in semi-arid and arid climates, the storms are more violent and a large amount of water falls in a short time. Under these conditions no soil is able to absorb the rain as rapidly as it falls, and hence a high percentage of it runs off even on soils of a sandy texture. The so-called dry washes, arroyos, and gulches of the Great Plains and the West bear witness to the erosive power of these torrential rains falling on lands without protective cover.

Regions with a large annual rainfall are also subject to severe erosion. Such areas usually have mild climates, and the soil is frozen only a small part of the year if at all. Under such conditions there is much more opportunity for soil washing than there is where soils are frozen for

several months each year. High rainfall in areas of rolling topography offers ideal conditions for erosion.

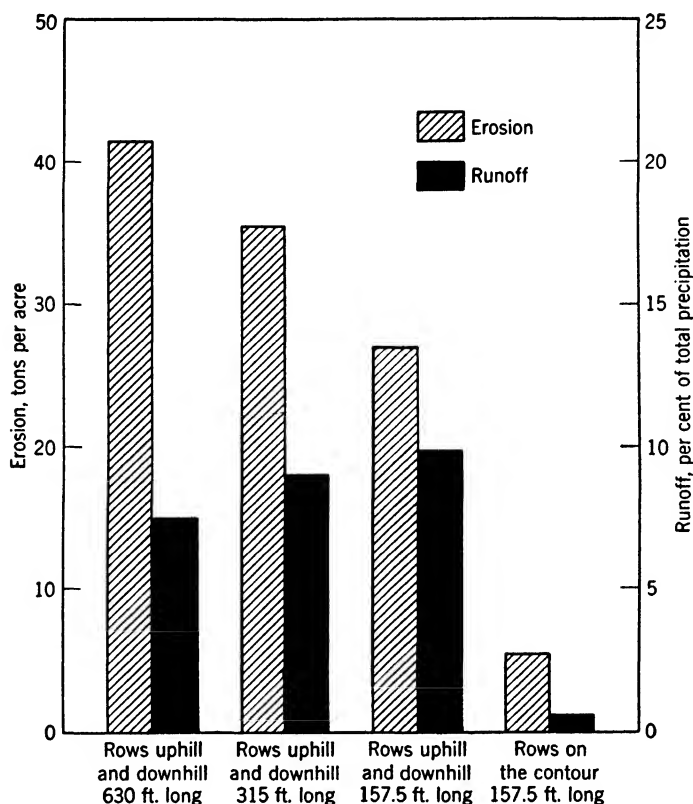


FIG. 81. Effect of length of slope on per cent of runoff water and loss of soil by erosion. Also a comparison of losses when corn is planted on the contour vs. up and down the slope. Soil was Marshall silt loam near Clarinda, Iowa. Annual precipitation 28.33 inches. Cropping system consisted of corn annually planted in lister furrows. Although these data show a direct relationship between length of slope and soil and water losses, Professor Krusekopf of Missouri found that this relationship did not hold when a rotation of corn, wheat, and clover was followed. [Figure from U.S.D.A. Tech. Bull. 959.]

**Degree and Length of Slope Pertinent Factors.** Other factors being equal, the steeper the slope the more rapidly will water run off over it. A fact to remember in this connection, however, is that the carrying capacity of water increases very rapidly with increase in veloc-

ity. For example, if the rate of flow is doubled, the water can carry particles sixty-four times the size of those carried previously. The quantity of soil carried by water running rapidly is also much greater than that carried by slowly moving water. In this connection one must remember that soil cannot be carried away until it is dislodged. Clear

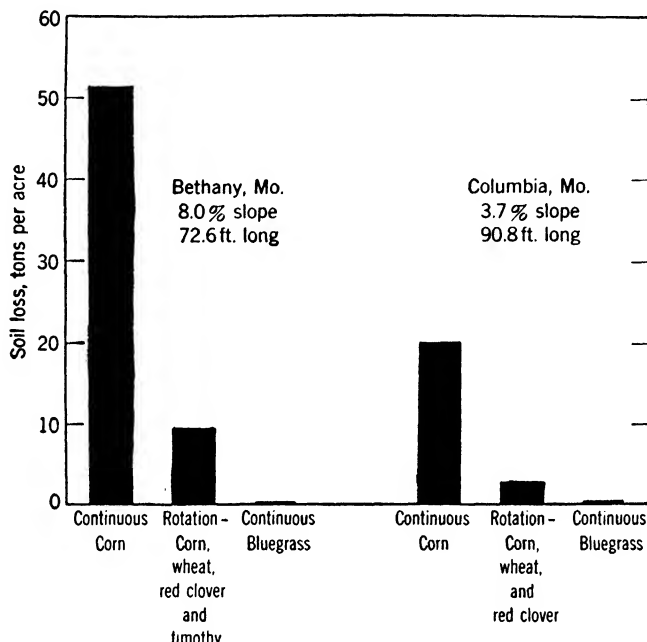


FIG. 82. Effect of steepness of slope and cropping system on loss of soil by erosion. Although the two slopes studied are not equal in length and the two fields are not at the same location, it is believed that conditions are sufficiently similar so that the data are comparable. [Missouri Agr. Exp. Sta. Bull. 518.]

water running over a soil surface at even a relatively high velocity was shown by Ellison to have little erosive power because it dislodged only a small amount of soil. When the water contained abrasive particles, however, erosion resulted because these particles cut loose the soil particles and then they could be transported by the water. It is readily seen then that steep slopes are more subject to erosion than gentle ones.

Long slopes of moderate steepness may suffer more erosion than steep but very short slopes. In fact, control of erosion on long moderate slopes is a greater problem than it is on short, steep slopes because it is

more difficult to find means for safely conducting a large volume of water over a long distance than it is to control the movement of a small volume a short distance even though the flow is more rapid.

**Time an Important Factor in the Absorption of Water by Soil.** As has been pointed out, a highly granular soil absorbs water much more readily than one which is in a single-grained or puddled structural condition. Granulation and stability of granules can be increased by growth of sod crops, good cultural practices, mulch covers, and increased organic matter content. Duley has shown that from two to four times as much water may run off of bare soil during a heavy rain as from soil protected by crop residues. Under most conditions the movement of water downward into a soil is a relatively slow process. As a result, many storms supply water at a faster rate than most soils can absorb it. It becomes necessary, therefore, to keep the soil permeable and to devise methods of slowing the movement of water over the soil surface, that is, *to reduce the rate of runoff* in order to provide time for a more complete absorption of it. This is one of the fundamental principles underlying erosion control practices.

**Use of Thick-Growing and Sod Crops to Decrease Erosion.** Crops which cover the ground surface and fill the surface soil with fibrous roots tend to hold it in place and reduce erosion. Dense sods which are produced by many grass species are outstanding in this respect. They are followed closely by several legumes such as lespedeza, crimson and various other clovers, and alfalfa, which develop a sod-like root formation in the surface soil. Other crops, such as kudzu, which rapidly cover the soil surface with a dense growth are also effective in reducing erosion. A vegetative cover prevents the direct impact of raindrops on the soil.

The small grains, oats, wheat, and barley, are much less efficient than sod-producing plants in protecting the soil surface from rain and controlling soil washing, but in turn they protect the soil to a far greater extent than do crops grown in cultivated rows.

Aside from the effect of the root system and the interception of falling rain, "close-growing" crops or crops which have many stems to the square foot decrease erosion by slowing down the movement of water over the soil surface. Each stem offers an obstruction to water movement, and when the stems are close together the rate of runoff is greatly reduced. This action not only reduces the erosive power of the water but also provides time for more of it to soak into the soil.

In planning a rotation for land which is subject to erosion liberal use should be made of hay, pasture, and small grain crops. The greater

the tendency toward erosion, the smaller is the proportion of time that cultivated crops should occupy the land. The data in Table 41 show the effect of different crops on the quantity of soil eroded.

TABLE 41

TONS OF SOIL LOST THROUGH EROSION PER ACRE PER YEAR AND PERCENTAGE OF PRECIPITATION WHICH RAN OFF UNDER DIFFERENT CULTURAL OR CROPPING SYSTEMS \*

(Average of 14 years. Slope 3.68%. Average precipitation 40.37 inches.)

| Cultural or Cropping System                    | Plowed<br>8 Inches<br>Uncropped | Rotation:        |                    |                      |                       |
|--|---------------------------------|------------------|--------------------|----------------------|-----------------------|
|  |                                 | Bluegrass<br>Sod | Wheat<br>Each Year | Wheat, and<br>Clover | Corn,<br>Each<br>Year |
| Tons of soil                                   | 41.08                           | 0.34             | 10.10              | 2.78                 | 19.72                 |
| Number of years to erode<br>7 inches of soil   | 24                              | 3,043            | 100                | 368                  | 50                    |
| Percentage of runoff                           | 30.3                            | 12.0             | 23.3               | 13.8                 | 29.4                  |
| Pounds of soil lost for<br>each inch of runoff | 6,734                           | 142              | 2,149              | 993                  | 3,314                 |

\* Prepared from "The Influence of Systems of Cropping and Methods of Culture on Surface Runoff and Soil Erosion," M. F. Miller and H. H. Krusekopf, *Missouri Exp. Sta. Res. Bull.* 177, 1932, p. 22.

**The Capability Factor in Land Management.** Cropping systems may be devised for soils on the basis of any one of the following characteristics: slope, erodibility, need for drainage, moisture-retaining capacity, depth, fertility. Any such plan is faulty because it takes into consideration only one soil characteristic, whereas in reality the performance of land when being cropped is the resultant of the interaction of all the soil properties and the environmental factors. It is more logical, therefore, to classify or group soils on the basis of the result which may be expected from a crop or cropping system. Under such a scheme the results from a given crop will be approximately the same on all soils in the same class, although the soil characteristics and environmental factors which bring about the result may vary considerably.

The Soil Conservation Service usually divides land into two general groups: (1) that suitable for cultivation, and (2) that not suitable for cultivation but suitable for permanent vegetative cover. Each of these groups is then divided into four classes on the basis of the degree of hazard or limitation encountered in their management. Classes I, II, and III include land which is suitable for regular cultivation. Land in class I can be farmed with the least danger of deterioration. Land in

class II will require some special management practices such as contouring or water control measures. Land in class III will require more extreme management, possibly terracing or the disposition of water on low wet areas. Land in class IV must be cultivated with great caution. It is usually better to leave this land in hay or pasture.

Classes V, VI, and VII include land not suited for cultivation. It should be used for grazing or forestry. In fact, land in class VI will need special care even for such uses because of steep slopes, thinness of soil, and similar characteristics. Land in class VII requires still more protective measures if it is to be used successfully for grazing or forestry. Class VIII includes land which is suited only for wildlife, recreation, and watershed protection. Its limitations may be extreme wetness, stoniness, steepness, erodibility.

Soils within a class may be further divided on the basis of the kind of hazards involved in their management. For example, one area may be subject to water erosion, while another area may be limited in use by a low moisture capacity. These subclasses may be divided again into land-capability units. This subdivision places in one unit those areas which require similar treatment.

The outline of the classification scheme is shown in Chart 8, and a land-capability map is presented in Fig. 83.

*Fitting the Crop to the Soil.* The growth habits of crops should be considered in selecting the crop to plant on different fields. For example, crops planted in rows and cultivated should be grown on the more level fields in so far as that is feasible, because erosion is excessive under such crops. On the other hand, hay and pasture crops can be more safely grown and are well adapted for growth on the more rolling fields. Small grains may be fitted into rotation on both relatively level and quite rolling land. Rotations on strongly sloping land should be long, with a number of years of hay or pasture and not over one year of grain crops. The steeper slopes may well be in pasture permanently or in wood lots. A little careful planning in regard to the crops to plant on different fields may well result in an increased production of cash and feed crops in addition to a better control of erosion.

**Strip Cropping a Practical Means of Reducing Erosion.** In view of the variation in the effect of different crops on erosion, it is logical to plant crops with different growth habits in strips across sloping fields. By this system erosion which may start in a strip of cultivated crop such as corn, cotton, or beans can be arrested in a strip of hay growing next down the slope. A strip of small grain can then follow, then another strip of hay, and then a cultivated crop. The width of

the strips of each crop should be determined by the steepness of the slope, the amount and nature of the precipitation, and the erodibility

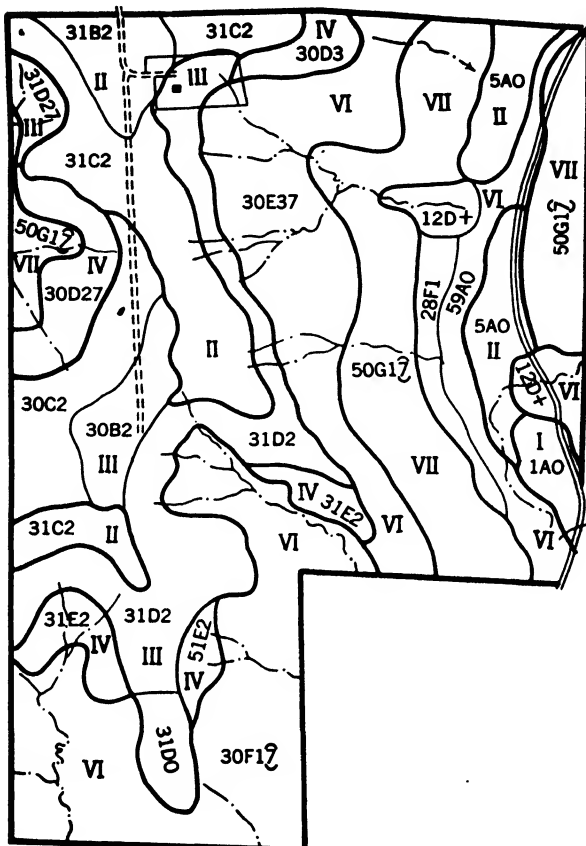


FIG. 83. Land-capability map of a 200-acre dairy farm in Wisconsin. The Roman numerals designate the land-capability classes. The other numbers and letters indicate the land characteristics. For example: in 30E37, the 30 designates the type of soil, the E indicates the steepness of slope, and the 37 stipulates that over 75 per cent of the topsoil has been lost and that there are occasional gullies. For explanation of land-capability classes see Chart 8. [Courtesy of Soil Conservation Service.]

of the soil. The strips should be planted on the contour as nearly as feasible.

At first thought the planting of crops in strips across a slope seems inconvenient and inefficient to many farmers. Experience, however,

Chart 8. Outline of the Land-Capability Classification

| Major Land-Use Suitability<br>(Broad grouping of limitations) | Land-Capability Class<br>(Degree of limitations)   | Land-Capability Subclass<br>(Kind of limitations)<br>(Groupings of land-capability units)<br>(Examples of possible subclasses<br>in class III)  | Land-Capability Unit<br>(Distinctive physical characteristics)<br>(Land-management groups based on<br>permanent physical factors)<br>(Examples) |
|---|--|---|---|
| Suited for cultivation  | I. Few limitations; wide latitude for each use; very good land from every standpoint<br><br>II. Moderate limitations or risks of damage; good land from all-round standpoint<br><br>III. Severe limitations or risks of damage; regular cultivation possible if limitations are observed<br><br>IV. Very severe limitations; suited for occasional cultivation or for some kind of limited cultivation<br><br>V. Not suited for cultivation; suited for permanent vegetation | I. Limited by hazard of water erosion; moderately sloping land<br>2. Limited by excess water; needs drainage for cultivation<br>3. Limited by low moisture capacity; sandy land<br>4. Limited by tight, very slowly permeable subsoils; claypan land  | 13C2. Moderately sloping, slightly acid soils on limestone<br>9C2. Moderately sloping, highly acid soils on sandstone or shale                  |
|   |  | V. Not suited for cultivation because of wetness, stones, overflows, etc.; few land limitations affect grazing or forestry use<br>VI. Too steep, stony, arid, wet, etc., for cultivation; moderate limitations for grazing or forestry<br>VII. Very steep, rough, arid, wet, etc.; severe limitations for grazing or forestry<br>VIII. Extremely rough, arid, swampy, etc.; not suited for cultivation, grazing, or forestry; suited for wildlife, water-sheds, or recreation | Land-management groups based on permanent physical factors such as range sites or forest sites  |

has proved it otherwise. Many times fences must be removed and a new field arrangement worked out. Strip cropping frequently results in longer rows and, hence, fewer turns with planting, tillage, and harvesting machinery. Also, since the strips follow the contour of the land, work is done approximately on the level in place of up- and downhill. This arrangement requires less power.



FIG. 84. Strip cropping, alone or in conjunction with terracing, makes possible the utilization of much land which would otherwise be severely damaged by erosion. [Courtesy of Soil Conservation Service.]

Strip cropping is adapted to land with long and fairly regular slopes. With a "choppy" topography involving short and very irregular slopes, such as is found in some glaciated areas, the system has less general application.

**Use of Terraces Increasing.** Many agriculturists in the northern states are inclined to look upon terracing as "something new" in soil management in America, but terracing is not new. Various types of ridges and depressions or channels across slopes were used in Colonial times to control runoff water. The term *terrace* was applied to these structures as early as 1847. From the Southeastern States the practice of terracing spread westward as farmers migrated in that direction. Terraces were used in Texas in 1903 and in Oklahoma by 1907. Use of

this type of erosion control is gradually developing in the northern states.

There are several types of terraces. Three types will be mentioned, but a detailed description of them and their uses cannot be given here. The bench terrace is the oldest type, having been used by ancient peoples in South America, Europe, and Asia. It is used primarily on steep slopes and where land for crop production is limited. The narrow, quite level benches are supported by very steep and sometimes practically vertical risers, sometimes covered with vegetation and sometimes consisting of retaining walls of rock. In the United States such terraces have been used to some extent in parts of the Southeastern States but are no longer recommended. They are still in use in certain citrus and avocado districts of California.

During the period immediately before and somewhat after 1800, farmers in southeastern United States began making shallow ditches across the slopes in their fields. These were known as "hillside ditches" and have developed into what is now known as *graded-channel* or *interception-and-diversion* terraces.

The most widely used type of terrace was developed around 1885 by a farmer named Priestly Mangum, near Wake Forest, North Carolina. This structure is known as the *broad-base* or Mangum terrace. It consists of a low, broad mound crossing the slope approximately on the contour. The shallow depression on the uphill side has sufficient fall to carry excess water around the hill to an outlet channel which carries it safely down the slope. The ridges may be constructed by repeated plowing of a strip of land, the soil always being thrown toward the center, or by plowing a strip and working the soil toward the center with a grader, whirlwind terracer, or one-way disk plow. No special equipment is needed for the construction of terraces. The ridges are of such low relief that farm implements may work over them with comparative ease. The entire field, including the ridges and depressions above them, may be planted to the crop to be grown. Crops should be planted on the contour.

**Sod Waterways Effective and Not Inconvenient.** Because no system of soil management will result in the complete absorption of all water from rain and melting snow, some provision must be made for conducting water down slopes with a minimum of gully erosion. Depressions or shallow channels into which water naturally drains as it moves down the slope may be kept in sod and so reduce gully erosion to a minimum. At times some grading may be necessary to bring the

water from various parts of a field into such a channel, and the drainageway itself may need to be broadened or the sides graded down to improve its usefulness. Some sodding is frequently necessary after alterations in the drainageway to prevent soil washing while a grassed waterway is being established by seeding. A straw mulch held down



FIG. 85. This grassed waterway is furnishing considerable hay in addition to permitting water to flow down the slope without damage to the soil. [Courtesy of Soil Conservation Service.]

by wire netting or brush is often useful in controlling erosion while a stand of grass is developing. On larger areas it is more practical to apply a straw mulch on a well-prepared seedbed and then work it in with a rotary treader so that part of the straws stand on end and protrude above the surface. When working in a field with sod waterways, the farmer lifts his tillage implements or straightens his disks in crossing the channel so as not to damage the sod. Sod waterways are a simple and effective device for preventing gully formation and for stopping the enlargement of small gullies. A large field may have several of them, some quite short and others of considerable length.

**CONTROLLING EROSION CAUSED BY WIND**

Wind erosion occurs on soil unprotected or only partially covered by vegetation. Soil blowing then may be expected during periods when land is being fitted for planting, before a young crop has reached sufficient size to protect the soil, and when land is being fallowed. Much wind erosion also occurs on range land which has been overgrazed.

Wind erosion reaches its greatest proportions in semi-arid and arid regions. Nevertheless, much damage is caused to both crops and soils in humid areas by soil blowing, although the phenomenon is less spectacular and attracts comparatively little attention in these regions. In the United States more attention has been given to the control of wind erosion in the Great Plains than in other sections of the country. In North Dakota damage from this cause was reported as early as 1888, and in Oklahoma four years after breaking of the sod. Soil blowing was one of the hazards confronting the early settlers, and control of wind erosion was one of the first problems studied by Agricultural Experiment Stations in the Plains states. The questions listed will call attention to some of the causes of wind erosion and some of the practices followed to control it.

**Questions**

1. What kinds of soil and soil particles blow?
2. Are losses from wind erosion confined to crop and soil damage?
3. Is there more than one type of soil blowing?
4. Should all soil be cultivated?
5. What farming practices lead to soil blowing, and which help control it?
6. How may seeding and planting methods be modified to reduce erosion?
7. How can rotations and strip cropping be used to reduce wind erosion?
8. What is meant by stubble-mulch culture?
9. What kinds of plants can be used to stabilize blowing sand?
10. What methods are used to control blowing on organic soils?

**Blowing Not Confined to Small Particles.** Sand grains and clusters or granules of small particles are not picked up by wind so easily nor carried such great distances as are silt and clay particles. Nevertheless, these larger particles are moved over the soil surface by rolling and by short skips. In this way they may do much damage to a young crop and may be piled up into small ridges and great dunes. Smaller particles may be carried along somewhat above the soil surface, or they may be swirled high in the air and carried long distances. Rain containing considerable fine soil, which was gathered from the air, was reported in

some of the eastern states during the great dust storms in the Plains states in the early 1930's.

**Blowing Not Confined to Sandy Soil.** When the prairie sod in the "short grass" or western Great Plains is plowed, the organic matter appears to decay rapidly during the first years of cultivation. One of the binding materials which holds the soil particles in clusters is thus destroyed. Furthermore, during periods of continued drought soil granules disintegrate, leaving the soil surface covered with a powdery layer which blows with extreme ease. It is seen, then, that, although sandy, incoherent soils offer less resistance to blowing when wind velocity is sufficiently high, loam and clay soils also blow when conditions induce disintegration of the soil granules and small clods.

Soils composed largely of the partially decayed remains of plants, variously called muck and peat in different states, have very low specific gravities. As a result, these soils blow readily during periods of dry weather when the surface is not protected by a vegetative cover. The fact that such soils constitute level areas adds to the likelihood of wind erosion.

**Crops Damaged by Blowing Soil.** If blowing starts while crops are small, the soil may be blown away from the root systems and the plants actually "blown out" over considerable areas. This is one type of damage greatly feared by onion growers on muck soil during dry springs. In dry falls, winter wheat may be blown out over large areas in the hard winter-wheat region.

Young crops may also suffer severe damage through being bruised by blowing soil particles. There may not be sufficient blowing to raise large clouds of dust, but the soil particles moving over or just above the surface cut and bruise the tender young plants. This type of damage is not uncommon on level sandy soils or on muck soils.

Another source of danger to crops is the possibility of being covered with blowing soil. A crop in a well-managed field may be covered by blowing soil from a poorly managed field near-by. Likewise, soil from higher land may be deposited on the crop growing in lower, more sheltered areas.

**Damage to Fences, Buildings, and Highways.** When blowing starts on a neglected or abandoned area of land, the soil will pile up around buildings and fences in an unbelievably short time.

Sand dunes around large lakes and in desert areas may cause much damage by moving over highways and encroaching on resort property. Summer cottages along lakes sometimes have to be protected from blowing sand.

**Three Types of Soil Blowing.** Soil particles move in three ways during wind erosion. Very fine soil particles, those from 0.1 to 0.15 mm. in diameter, are rolled over the surface by direct wind pressure and then suddenly jump up almost vertically from a short distance to a foot or more. Once in the air, the particles gain velocity and then descend in an almost straight line, not varying more than 6 to 12 degrees from the horizontal. The horizontal distance traveled by a particle is from 4 to 5 times the height of its jump. Upon striking the surface, the particles may rebound into the air or knock other particles into the air and come to rest themselves. The major part of the soil carried by wind moves by this process, which is called *saltation*. It is interesting to note that around 93 per cent of the total soil movement by wind takes place below a height of 1 foot, and probably 50 per cent or more occurs between 0 and 2 inches.

Very fine dust particles are protected from wind action because they are too small to protrude above a minute viscous layer of air which clings to the soil surface. As a result, a soil composed entirely of extremely fine particles is very resistant to wind erosion. These dust particles are thrown into the air chiefly by the impact of particles moving in saltation, but once in the air their movement is governed by wind action. They may be carried very high and over long distances.

Relatively large particles (between 0.5 and 1.0 mm. in diameter) are too heavy to be lifted by wind action but are rolled or pushed along the surface by the impact of particles in saltation. This process is called *surface creep*.

From the above facts it is evident that wind erosion is due principally to the effect of wind on particles of a suitable size to move in saltation. Accordingly, wind erosion can be controlled if (1) the soil particles can be built up into clusters or granules of too large a size to move in saltation; (2) the wind velocity near the soil surface can be reduced by ridging the land, by vegetative cover, or even by developing a cloddy surface; and (3) by providing strips of stubble or other vegetative cover sufficient to catch and hold the particles moving in saltation. Some management practices designed to provide these conditions are discussed in the following paragraphs.

**Soil Areas To Be Left in Sod.** Studies have shown that extensive wind erosion originates mainly in land where crops frequently fail because of inadequate moisture supply. Sandy soils and shallow soils suffer the most, particularly in the areas receiving 14 to 17 inches of annual rainfall. Accordingly, land in regions of low rainfall, where production of a crop is dependent on a "favorable" season, is more

safely left in native grasses. Unfortunately, during periods of high demand for food with correspondingly high prices, large areas of such land have been plowed and planted. Serious soil blowing is certain to result, and very quickly when a series of normal or of unusually dry years follows. Furthermore, it is very difficult to reestablish the sod on such land when wind erosion has once gotten well under way.

**Summer Fallow Dangerous.** Any cropping system which leaves the land unprotected by a growing crop for an appreciable length of time opens the opportunity for soil blowing. In regions where annual precipitation or precipitation during the growing period is insufficient to produce a crop, either some system of moisture storage in the soil or irrigation must be resorted to. Keeping the soil in a fallow condition with enough cultivation to prevent weed growth is a practice frequently followed under such conditions. Unfortunately, there has been considerable trouble with soil blowing during the fallow year in parts of the Great Plains. Alternating strips of a summer crop such as sorghum with strips of fallow has proved an effective way of reducing soil blowing when fallowing is necessary. The use of subsurface tillage, as discussed in the section on stubble-mulch culture below, also offers a satisfactory means of handling fallow ground.

**Pulverization of Soil Not Desirable.** In cultivating a soil being maintained in a fallow condition and in fitting seedbeds for any crop in areas subject to wind erosion, the soil should never be pulverized. The objective should be to leave the soil in a cloddy condition. A cloddy surface soil contains a minimum of particles of the size which blows readily, and, furthermore, the clods reduce wind velocity near the soil surface. In view of these facts it can be seen that rollers, harrows, and other implements which tend to make the soil fine have little use where soil blowing is a hazard. Some agronomists suggest that the moldboard plow be used sparingly in such areas. The lister and the duckfoot or sweep-type cultivator are two implements used widely in areas subject to wind erosion.

**A Smooth Soil Surface Undesirable.** Furrows made by the lister or duckfoot cultivator at right angles to the wind are helpful in reducing soil blowing. When these furrows become filled with soil, new ones should be made.

Another method of keeping the soil surface rough is to plant crops in furrows instead of on the level, as is done in humid regions. Row crops such as corn and members of the sorghum family are planted in lister furrows. Drills which make shallow furrows have also been developed

for planting small grains. Naturally all plantings should be made at right angles to the prevailing winds.

**Rotations Helpful.** Rotations which involve a minimum of soil tillage are helpful in preventing wind erosion. A typical rotation consists of one year of cultivated crop, then a spring-seeded crop, a year



FIG. 86. Listing on the contour not only protects the listed field from wind action but also may collect much soil blowing from an adjoining field, as illustrated in this Oklahoma field. The velocity of the wind is indicated by the posture of the man in the foreground. [Courtesy of Soil Conservation Service.]

of fallow, followed by winter wheat. Alfalfa makes a wind-resistant sod but is objectionable in areas of low rainfall because it dries out the soil to a considerable depth. A rotation suggested by Professor Nelson of the Wyoming Agricultural Experiment Station consists of a cultivated crop followed by a spring-drilled crop, then rye, a fallow year, and winter wheat. Some wheat farmers object to rye because it becomes mixed with the wheat and lowers the market grade. Land in grass is maintained for a long time because of the difficulty of establishing a sod. Other rotations or sequences of crops suitable for growth in a given locality and arranged in an order to meet the hazards of wind and water

erosion on given farms or fields can be obtained from state colleges and local technicians of the Soil Conservation Service.

**Strip Cropping Effective.** The planting of crops in strips at right angles to the prevailing winds is proving valuable in reducing soil blowing. The width of the strips is determined by the nature of the soil, exposure to wind, and similar factors. On sandy soils the strips may be



FIG. 87. By working crop residues partially into the soil, leaving a considerable portion on top, this Montana soil is protected from wind erosion. The soil is also in an excellent condition to absorb rainfall. The work is being done with a one-way plow. [Courtesy of Soil Conservation Service.]

only 8 to 10 rods wide, but on heavier soils they may be from 15 to 20 rods in width.

**Stubble-Mulch Culture Helpful.** Crop residues are an effective soil protection against wind erosion. Small-grain stubble reduces wind velocity and also catches soil particles moving in saltation. Strips of stubble left at frequent intervals across a field being fallowed or fitted for a spring crop form effective barriers. The height of the stubble as well as wind velocity influence the width of strip needed, but a 10-foot strip of short stubble should trap 90 to 98 per cent of the moving soil.

Another effective cultural practice is to work crop residues partially into the soil but leave enough on the surface to obstruct the movement

of soil particles and reduce the force of the wind. The one-way disk and various types of subsurface tillers are useful in this type of soil culture. Following the binder or combine with a subtiller with a "treader" hitched behind it is a good way to leave soil in a moisture-receptive condition and also protected from wind and water erosion. A treader is a type of rotary hoe with the pronged wheels reversed. A treader is also a useful implement for killing weeds and smoothing and packing subtilled ground in preparation for seeding.

In planting crops in the mulch-culture system of farming, drills and planters must be used that will work through the crop residues without covering them or leaving them in such a position as to interfere with seed germination or growth of young plants. The semi-deep-furrow drill or the deep-furrow drill accomplishes these purposes in seeding small grains and puts the seed down in moist soil. The low-down, press-wheel drill also is satisfactory. Row crops may be planted with a standard planter having furrow openers mounted on stub runners which pass through the residues with little trouble.

Crops should be cultivated in such a way as to leave the residues on the surface; in other words, cultivate beneath them. This result can be accomplished by use of sweep-type cultivators. The stubble-mulch system of soil management has many possibilities and should be carefully investigated in areas where wind or water erosion is a serious hazard.

**Stabilization of Blowing Sand with Trees and Coarse Grasses.** Sand dunes and hilly sandy areas which are blowing badly in humid areas are effectively stabilized by use of coarse grass, such as one of the beach grasses, and by reforestation. The grasses are transplanted in little bunches. Originally these were placed in a double row with plants about 18 inches apart. The pairs of rows were spaced from 12 to 40 feet according to conditions. Later, planting the pairs of rows in a serpentine pattern rather than straight was considered more effective. A more recent system is to plant the rows in a radiating pattern, like the spokes of a wheel, or in the form of a cross. This method of planting requires far less material and labor than the older systems.

The grass spreads by means of rhizomes and will establish a thick enough sod to hold the sand. Sometimes it is necessary to protect the new plantings by windbreaks of fencing or brush. Trees or shrubs are often planted to supplement the effect of the grass. With the older and thicker systems of setting the grass, trees may be planted immediately, but with the newer systems several years should elapse before the trees are planted.

The establishment of a forest cover on blowing sand plains and hills is desirable but is sometimes beset with difficulties. The sand may be blown out from around the young trees, or they may be severely cut by the blowing sand or covered with it unless some protective measures are adopted. Temporary windbreaks afford protection, and in special small areas the soil may be covered with a mulch of straw which is held in place by brush. This practice, however, is expensive.

In humid sections various species of conifers have proved satisfactory for planting on sandy soils subject to blowing, and in some areas the black locust has been used very successfully. Trees suitable for use under the climatic conditions which prevail in any given area should be selected.

In drier areas a coarse-growing plant such as sorghum or sudan grass may be grown and the residues worked into the soil to control blowing while a grass is being established.

**Trees for Windbreaks on Organic Soils.** Organic soil areas of appreciable size are frequently protected from wind damage by planting windbreaks of trees around them. In addition, rows of trees are often planted across the area at right angles to the prevailing wind. Willows have been found satisfactory for this purpose, especially since they make a very rapid growth in organic soil. A number of species of conifers have also been used. Austrian and Scotch pine make a rapid growth and give promise of making satisfactory windbreaks in the North Central states. White pine grows more slowly but otherwise appears suitable for such use. The use of trees as windbreaks must be limited because of the large amount of soil they take out of crop production on account of their extensive root system. A number of shrubs also make good windbreaks. Spirea is frequently used.

**Additional Protection from Wind Needed.** Because of the limitation in use of trees, supplementary windbreaks are needed on muck areas of appreciable size. Picket fencing, such as is used for keeping the snow off of highways in the northern states, makes very satisfactory temporary windbreaks. Fastening burlap bags to the lower part of the fence increases its effectiveness. Bags also may be hung from wires stretched between posts driven into the muck.

Occasionally rows of corn have been planted across the field at frequent intervals. A more common method is to drill five or six rows of rye across the field at intervals of 50 or more feet. All such plantings must be at right angles to the wind. Planting adapted grasses on ditch banks has been found a satisfactory method of protecting these exposed areas in Michigan and preventing filling of the ditch with wind-carried muck.

Moist muck does not blow appreciably. Accordingly, use of an overhead irrigation system is helpful during windy periods before the crop cover is sufficient to prevent soil blowing. Rolling the muck with a very heavy roller induces moisture to rise more rapidly by capillarity and so dampens the surface layer. This practice is not effective unless the soil layers below the surface are quite moist.

# 17

## Soil Fertility Maintenance and Productivity Rating of Soil

Differences in the productive capacity of different kinds of soils, or land, are commonly recognized on the basis of experience in farming. The range in productive capacity of different soil types is very evident where the same crops are grown and the same general farming practices are followed; but, even where practices are modified and crop adaptation is recognized, differences in yields, or net money income, still prevail on widely different soil types. Some system of expressing the difference in productivity of soil types or land areas is highly desirable. Furthermore, the importance of maintaining soil productivity, to farmers, to urban workers, and to the nation, is well recognized and is stressed in another section of this book. But the question may well be asked, how does a farmer know whether or not he is maintaining the productive capacity of his soil? It might be assumed that, if crop yields do not decrease, fertility is being maintained. This is a false assumption, however, for the use of better tillage methods, of more carefully selected seeds and seeds of improved crop varieties, and of more suitable rates, dates, and methods of planting may maintain or even increase yields for a period in spite of a lower soil fertility. The discussion of the problems of measuring fertility maintenance and of rating the productivity of land is the objective of this chapter.

### Objectives

- A. Maintaining soil fertility.
- B. Soil-productivity rating and land classification.

### MAINTAINING SOIL FERTILITY

In recent years some system has been sought to determine if a given rotation or soil-management system is maintaining the productivity

of the soil and, if not, how rapidly fertility is being lost. Workers at the experiment stations in Ohio and Missouri have taken the lead in the endeavor, and the systems proposed by them are discussed in connection with the following questions.

### Questions

1. What productivity indexes are given to different groups of crops?
2. On what basis are productivity indexes determined?
3. Why is so much emphasis placed on nitrogen in assigning index values to crops?
4. Of how much value is manure in maintaining production?
5. Do fertilizers contribute to increased productivity?
6. How detrimental is erosion?

**Crop Indexes.** The Ohio and Missouri systems give to each crop a productivity index based on the effect of growing that crop on the nitrogen or organic matter content of the soil. Crops grown in rows and intertilled are given the highest minus ratings, whereas leguminous crops which have a sod-like habit of growth are given the highest positive ratings. The actual soil-improving values assigned to legumes depends on the age of the stand and the disposition of the crop. As would be expected, the highest value results from returning the entire crop to the soil, and the lowest when a hay crop is harvested.

Grass sods are given a zero rating unless the crop is grazed off or harvested for hay, in which case a rating of -0.50 per cent is given. Grasses add no nitrogen or mineral nutrients to the soil. They do, however, reduce erosion and loss of nutrients through leaching. Moreover, the organic content of the soil is not depleted through tillage. The growth of these crops should also encourage non-symbiotic nitrogen fixation. When a legume is grown in association with a grass, a positive index is given in accord with the proportion of the legume present. The indexes ascribed to different crops by different experiment stations vary slightly.

**Basis for Crop Indexes.** The productivity ratings assigned to different crops in Ohio were arrived at through a study of data from the Ohio Experiment Station. These data show that, when corn, oats, or wheat were grown year after year on the same land, the organic content of the soil was greatly decreased, especially by corn, and that the yields decreased correspondingly. When these crops were included in rotations with clover or alfalfa, the decrease in the nitrogen content of the soil was much less, as was also the decrease in crop yield; and, furthermore, the greater the number of years that clover or alfalfa occupied

the soil in the rotation period, the more effective was the rotation in maintaining soil productivity. A rotation in Ohio, including 3 years of alfalfa and 1 each of corn and oats, showed no decrease in corn yield for a 20-year period and no loss of soil organic matter. On the other hand, a rotation consisting of 3 years of corn, 1 of clover, and 1 of wheat, resulted in a decrease in corn yield of 21.6 per cent in 20 years and a loss of 18 per cent of the soil nitrogen.

Likewise, Missouri scientists base their crop indexes on results obtained from soil-management studies which have continued for many years. A direct relationship was found between the content of nitrogen in the plow soil of different soil types and the yield of corn. The results are presented in Table 42. The productivity indexes assigned to various crops handled in different ways are given in Table 43.

TABLE 42

## THE RELATION OF TOTAL NITROGEN CONTENT OF SOILS TO CORN YIELDS \*

| Soil Type          | Total Nitrogen<br>(pounds per<br>acre, 7 inches<br>deep) | Average Yield<br>of Corn<br>(bushels per<br>acre) | Pounds of<br>Soil Nitrogen<br>per Bushel<br>of Corn |
|--------------------|--|---|---|
| Marshall silt loam | 3,630  | 38.6  | 94.04   |
| Grundy silt loam   | 3,370  | 32.0  | 105.31  |
| Eldon silt loam    | 3,160  | 31.2  | 101.28  |
| Crawford silt loam | 2,840  | 25.4  | 111.81  |
| Cherokee silt loam | 1,950  | 22.5  | 86.67   |
| Gerald silt loam   | 1,890  | 19.0  | 99.47   |
| Union silt loam    | 1,600  | 16.0  | 100.00  |
| Average            | 2,634  | 26.4  | 99.92   |

\* "Evaluating Annual Changes in Soil Productivity," A. W. Klemme and O. T. Coleman, *Missouri Exp. Sta. Bull.* 405, 1939, p. 4.

**Nitrogen—the Key Element.** The question may well be raised, why is soil nitrogen content made the basis for the fertility ratings of crops instead of some other element, like phosphorus, required by plants? In this connection it must be remembered that nitrogen is accumulated in soils of humid regions almost entirely in the form of humus, which in turn is largely a product of plant-tissue decay together with the cells of microorganisms. In other words, to accumulate humus it is necessary for soil conditions to be favorable for a considerable if not luxuriant growth of plants, including legumes. This growth in turn involves at least a moderate supply of all essential plant-food elements, a not-un-

TABLE 43

PER CENT OF TOTAL NITROGEN GAINED OR LOST ANNUALLY FOR THE  
VARIOUS CROPS AND METHODS OF UTILIZING THEM \*

| Crops   | All Turned<br>Under or<br>Left on<br>Land | Grazed Off<br>or Only the<br>Grain or Seed<br>Removed | Whole Crop<br>Removed or<br>Burned |
|---|---|---|------------------------------------|
| (Productivity indexes given are for full stand and normal growth. One-half full crop means one-half the index listed, etc.)   |   |   |                                    |
| 1. Grasses  |   |   |                                    |
| Permanent pastures  | 0.00                                      | 0.00  | -0.50                              |
| Perennial grasses   | 0.00                                      | 0.00  | -0.50                              |
| Perennial grasses + 50% legumes   | +0.50                                     | +0.35   | 0.00                               |
| Sudan grass   | 0.00                                      | -0.50   | -2.00                              |
| 2. Intertilled crops  |   |   |                                    |
| Corn, tobacco, sorghums, etc. †   | 0.00                                      | -1.35   | -2.00                              |
| Cotton  | 0.00                                      | Lint &<br>seed -1.35                                  | -2.00                              |
| 3. Legumes—when well inoculated   |   |   |                                    |
| Alfalfa—over 1 year old   | +3.00                                     | +2.25   | +0.50                              |
| Alfalfa—under 1 year old  | +1.00                                     | +0.75   | 0.00                               |
| Clover, alsike—1st year   | +0.75                                     | +0.50   | 0.00                               |
| Clover, alsike—2nd year   | +1.75                                     | +1.30   | 0.00                               |
| Clover, crimson—spring after seeding  | +1.50                                     | +1.00   | 0.00                               |
| Clover, red—1st year  | +1.00                                     | +0.75   | 0.00                               |
| Clover, red—2nd year—1st crop   | +2.00                                     | +1.50   | +0.25                              |
| Clover, red—2nd year—2nd crop   | +0.50                                     | +0.35   | 0.00                               |
| Clover, sweet—1st year  | +1.00                                     | +0.75   | 0.00                               |
| Clover, sweet—2nd year  | +3.00                                     | +2.25   | +0.50                              |
| Clover, white or hop  | +1.00                                     | +0.75   | 0.00                               |
| Crotalaria  | +2.50                                     | +1.85   | 0.00                               |
| Lespedeza—crop growth to Aug. 1   | +1.00                                     | +0.75   | 0.00                               |
| Lespedeza—crop growth after Aug. 1  | +0.75                                     | +0.50   | 0.00                               |
| Lespedeza—total season crop   | +1.75                                     | +1.30   | 0.00                               |
| Soybeans, cowpeas, etc.   | +1.00                                     | +0.75   | -0.50                              |
| Vetch, winter   | +1.50                                     | +1.00   | -0.50                              |
| 4. Small grains   |   | Combined -0.65  |                                    |
| Wheat, oats, rye, barley, flax, etc.  | 0.00                                      | Grazed -0.25  | -1.00                              |
| 5. Soil treatments  |   |   |                                    |
| Barnyard or straw manure per ton applied  |   |   | +0.15                              |
| Commercial fertilizer—For each 40 lb. available plant food applied  |   |   | +0.15                              |
| All Crops turned under, grazed or removed from the land during the season should receive consideration.   |   |   |                                    |
| Heavy applications of straw or other organic matter low in protein without the growth of legumes on the land or the use of chemical treatments may cause a temporary reduction in available nitrogen.   |   |   |                                    |
| 6. Each ton of feed hauled to the farm and fed in the fields and not otherwise given credit as manure should be given positive credits as follows: small-grain straw or silage, 0.075; timothy, orchard grass, reedtop, sorghums, corn fodder, etc. (air dry), 0.225; legumes, grains, grain mixtures, and supplementary feeds, 0.45. |   |   |                                    |

\* From *Missouri Exp. Sta. Bull.* 522.

† Corn all hogged down should be given a factor of -0.50.

favorable soil reaction and drainage condition, and a reasonable amount of precipitation or irrigation water. On the whole, then, the nitrogen or humus content of a soil is a fairly accurate index of productivity.

**Value of Manure.** Manure contributes to soil productivity through both its humus content and the plant nutrients supplied which stimulate crop growth. The two contributions may be considered about equal in value in increasing the humus of the soil. The Missouri station determined the effect of a 6-ton annual application of manure for a period of 50 years on the nitrogen content of soil that was growing various sequences of crops. The total increase in nitrogen content of the soil was 41.1 per cent. As 300 tons of manure were applied, this makes an increase of 0.137 per cent of nitrogen for each ton of manure. A fertility index of 0.15 per cent is assigned for each ton of manure applied, as shown in Table 43.

**Fertilizer Application.** A study of the effect of applications of fertilizer on the nitrogen content of soil has led the Ohio station to give a positive index value of 0.15 per cent for each 200 pounds of fertilizer containing approximately 20 per cent of plant nutrients. The Missouri station arrived at a similar value.

**Detrimental Action of Erosion.** For many years the Missouri station has studied the loss of soil by erosion under different cropping systems on a field having a slope of 3.7 per cent. More recently many studies of erosion losses and of the effect of the loss of surface soil on productivity have been made by the Soil Conservation Service and by different experiment stations. A consideration of the available data has led to the compilation of a rather detailed table of factors to be applied in computing the detrimental effects of erosion. Length and per cent of slope are taken into consideration in association with the cropping system and the erosion control measures used. A portion of the table contained in *Missouri Experiment Station Bulletin 522* is presented in Table 44. The method of applying the soil-productivity factors or indexes of crops to a farm, together with deductions for erosion, and positive values for applications of manure and fertilizer are illustrated in Table 45.

#### SOIL-PRODUCTIVITY RATING AND LAND CLASSIFICATION

Generally, in the past, differences in the productive capacity or economic value of soils have been expressed loosely and qualitatively by such terms as "good," "fair," and "poor" or by the use of the economist's terms "supermarginal," "marginal," and "submarginal."

TABLE 44

EROSION FACTORS OR PER CENT OF SURFACE 7 INCHES OF SOIL LOST ANNUALLY FROM WATER EROSION FOR DIFFERENT CROPS OR CROPPING SYSTEMS ON DIFFERENT DEGREES AND LENGTHS OF SLOPE \*

| Crop and Sequence   | Slope Length (feet) | Crop or Rotation without Control Measures |     |     |      |      |      | Crop or Rotation plus Contouring |     |     |     |     |      | Crop or Rotation plus Contouring<br>Terraces and Contouring |     |     |     |     |     |
|---|---------------------|---|-----|-----|------|------|------|----------------------------------|-----|-----|-----|-----|------|---|-----|-----|-----|-----|-----|
|   |                     | 2   | 4   | 6   | 8    | 10   | 12   | 2                                | 4   | 6   | 8   | 10  | 12   | 2   | 4   | 6   | 8   | 10  | 12  |
| Intertilled crop; after small grain or <i>lespedeza</i> hay; or 1 year of intertilled following grass or legume crops | 200                 | 0.9                                       | 1.7 | 2.9 | 4.3  | 5.7  | 7.2  | 0.5                              | 0.9 | 1.5 | 2.3 | 3.2 | 4.5  | 0.2   | 0.6 | 1.1 | 1.8 | 2.6 | 3.6 |
|   | 400                 | 1.3                                       | 2.6 | 4.3 | 6.4  | 8.5  | 10.8 | 0.8                              | 1.3 | 2.2 | 3.4 | 4.8 | 6.8  | 0.2   | 0.6 | 1.1 | 1.8 | 2.6 | 3.6 |
|   | 600                 | 1.7                                       | 3.3 | 5.5 | 8.1  | 10.7 | 13.7 | 1.0                              | 1.7 | 2.8 | 4.3 | 6.1 | 8.6  | 0.2   | 0.6 | 1.1 | 1.8 | 2.6 | 3.6 |
|   | 800                 | 2.0                                       | 4.0 | 6.6 | 9.8  | 13.0 | 16.6 | 1.2                              | 2.0 | 3.4 | 5.2 | 7.4 | 14.4 | 0.2   | 0.6 | 1.1 | 1.8 | 2.6 | 3.6 |
|   | 1,000               | 2.3                                       | 4.5 | 7.5 | 11.1 | 14.7 | 18.8 | 1.4                              | 2.3 | 3.9 | 5.9 | 8.4 | 11.8 | 0.2   | 0.6 | 1.1 | 1.8 | 2.6 | 3.6 |
| Intertilled crop; after clover, green manure, or full-growth <i>lespedeza</i> turned under or worked in soil          | 200                 | 0.4                                       | 0.9 | 1.5 | 2.2  | 2.9  | 3.7  | 0.3                              | 0.4 | 0.7 | 1.1 | 1.6 | 2.3  | 0   | 0.1 | 0.4 | 0.8 | 1.2 | 1.7 |
|   | 400                 | 0.7                                       | 1.3 | 2.2 | 3.2  | 4.3  | 5.5  | 0.4                              | 0.7 | 1.1 | 1.7 | 2.4 | 3.4  | 0   | 0.1 | 0.4 | 0.8 | 1.2 | 1.7 |
|   | 600                 | 0.8                                       | 1.7 | 2.8 | 4.1  | 5.5  | 6.9  | 0.5                              | 0.8 | 1.4 | 2.2 | 3.1 | 4.3  | 0   | 0.1 | 0.4 | 0.8 | 1.2 | 1.7 |
|   | 800                 | 1.0                                       | 2.0 | 3.3 | 5.0  | 6.6  | 8.4  | 0.6                              | 1.0 | 1.7 | 2.6 | 3.8 | 5.3  | 0   | 0.1 | 0.4 | 0.8 | 1.2 | 1.7 |
|   | 1,000               | 1.1                                       | 2.3 | 3.8 | 5.6  | 7.4  | 9.5  | 0.7                              | 1.1 | 2.0 | 3.0 | 4.2 | 6.0  | 0   | 0.1 | 0.4 | 0.8 | 1.2 | 1.7 |
| Intertilled crop; after meadow or pasture in which grass predominates   | 200                 | 0.3                                       | 0.6 | 0.9 | 1.4  | 1.8  | 2.3  | 0.2                              | 0.3 | 0.5 | 0.7 | 1.0 | 1.5  | 0   | 0   | 0.1 | 0.4 | 0.6 | 1.0 |
|   | 400                 | 0.4                                       | 0.8 | 1.4 | 2.1  | 2.7  | 3.5  | 0.3                              | 0.4 | 0.7 | 1.1 | 1.6 | 2.2  | 0   | 0   | 0.1 | 0.4 | 0.6 | 1.0 |
|   | 600                 | 0.5                                       | 1.1 | 1.8 | 2.6  | 3.5  | 4.4  | 0.3                              | 0.5 | 0.9 | 1.4 | 2.0 | 2.8  | 0   | 0   | 0.1 | 0.4 | 0.6 | 1.0 |
|   | 800                 | 0.6                                       | 1.3 | 2.1 | 3.2  | 4.2  | 5.3  | 0.4                              | 0.6 | 1.1 | 1.7 | 2.4 | 3.4  | 0   | 0   | 0.1 | 0.4 | 0.6 | 1.0 |
|   | 1,000               | 0.7                                       | 1.5 | 2.4 | 3.6  | 4.7  | 6.0  | 0.4                              | 0.7 | 1.3 | 1.9 | 2.7 | 3.8  | 0   | 0   | 0.1 | 0.4 | 0.6 | 1.0 |

|  |              |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
|--|--------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Fall grain, without meadow seeding, after<br>intertilled, small-grain, or other annual<br>drilled crop; spring grain with meadow seed-<br>ing, after small grain or other annual drilled<br>crop | 200          | 0.4 | 0.8 | 1.4 | 2.1 | 2.7 | 3.5 | 0.3 | 0.4 | 0.7 | 1.1 | 2.2 | 0   | 0.1 | 0.4 | 0.7 | 1.1 | 1.6 |     |
|  | 400          | 0.6 | 1.3 | 2.1 | 3.1 | 4.1 | 5.2 | 0.4 | 0.6 | 1.1 | 1.6 | 2.3 | 3.3 | 0   | 0.1 | 0.4 | 0.7 | 1.1 | 1.6 |
|  | 600          | 0.8 | 1.6 | 2.6 | 3.9 | 5.2 | 6.6 | 0.5 | 0.8 | 1.4 | 2.1 | 3.0 | 4.1 | 0   | 0.1 | 0.4 | 0.7 | 1.1 | 1.6 |
|  | 800          | 1.0 | 1.9 | 3.2 | 4.7 | 6.3 | 8.0 | 0.6 | 1.0 | 1.6 | 2.5 | 3.6 | 5.0 | 0   | 0.1 | 0.4 | 0.7 | 1.1 | 1.6 |
|  | 1,000        | 1.1 | 2.2 | 3.6 | 5.4 | 7.1 | 9.1 | 0.7 | 1.1 | 1.9 | 2.8 | 4.0 | 5.7 | 0   | 0.1 | 0.4 | 0.7 | 1.1 | 1.6 |
| Spring grain with meadow seeding, after inter-<br>tilled crops; fall grain with meadow seeding,<br>after small grain or other annual drilled<br>crop   | 200          | 0.3 | 0.6 | 1.0 | 1.5 | 2.0 | 2.6 | 0.2 | 0.3 | 0.5 | 0.8 | 1.1 | 1.6 | 0   | 0   | 0.2 | 0.4 | 0.7 | 1.1 |
|  | 400          | 0.5 | 0.9 | 1.4 | 2.3 | 3.0 | 3.9 | 0.3 | 0.5 | 0.8 | 1.2 | 1.7 | 2.4 | 0   | 0   | 0.2 | 0.4 | 0.7 | 1.1 |
|  | 600          | 0.6 | 1.2 | 1.8 | 2.9 | 3.8 | 4.9 | 0.4 | 0.6 | 1.0 | 1.5 | 2.2 | 3.1 | 0   | 0   | 0.2 | 0.4 | 0.7 | 1.1 |
|  | 800          | 0.7 | 1.4 | 2.2 | 3.5 | 4.6 | 5.9 | 0.4 | 0.7 | 1.2 | 1.9 | 2.6 | 3.7 | 0   | 0   | 0.2 | 0.4 | 0.7 | 1.1 |
|  | 1,000        | 0.8 | 1.6 | 2.5 | 4.0 | 5.2 | 6.7 | 0.5 | 0.8 | 1.4 | 2.1 | 3.0 | 4.2 | 0   | 0   | 0.2 | 0.4 | 0.7 | 1.1 |
| Small grain with meadow seeding; after sod<br>crops  | 200          | 0.2 | 0.3 | 0.6 | 0.8 | 1.1 | 1.4 | 0.1 | 0.2 | 0.3 | 0.4 | 0.6 | 0.9 | 0   | 0   | 0   | 0.1 | 0.3 | 0.5 |
|  | 400          | 0.3 | 0.5 | 0.8 | 1.2 | 1.7 | 2.1 | 0.2 | 0.3 | 0.4 | 0.7 | 0.9 | 1.3 | 0   | 0   | 0   | 0.1 | 0.3 | 0.5 |
|  | 600          | 0.3 | 0.6 | 1.1 | 1.6 | 2.1 | 2.7 | 0.2 | 0.3 | 0.5 | 0.8 | 1.2 | 1.7 | 0   | 0   | 0   | 0.1 | 0.3 | 0.5 |
|  | 800          | 0.4 | 0.8 | 1.3 | 1.9 | 2.5 | 3.3 | 0.2 | 0.4 | 0.7 | 1.0 | 1.5 | 2.0 | 0   | 0   | 0   | 0.1 | 0.3 | 0.5 |
|  | 1,000        | 0.4 | 0.9 | 1.5 | 2.2 | 2.9 | 3.7 | 0.3 | 0.4 | 0.8 | 1.2 | 1.6 | 2.3 | 0   | 0   | 0   | 0.1 | 0.3 | 0.5 |
| Grass and legume meadows and pastures  | 200          | 0   | 0.1 | 0.2 | 0.2 | 0.3 | 0.4 | 0   | 0   | 0.1 | 0.2 | 0.2 | 0.2 | 0   | 0   | 0   | 0   | 0   | 0   |
|  | 400          | 0.1 | 0.1 | 0.2 | 0.4 | 0.5 | 0.6 | 0   | 0.1 | 0.1 | 0.2 | 0.3 | 0.4 | 0   | 0   | 0   | 0   | 0   | 0   |
|  | 600          | 0.1 | 0.2 | 0.3 | 0.4 | 0.6 | 0.8 | 0.1 | 0.1 | 0.2 | 0.3 | 0.5 | 0   | 0   | 0   | 0   | 0   | 0   | 0   |
|  | 800          | 0.1 | 0.2 | 0.4 | 0.5 | 0.7 | 0.9 | 0.1 | 0.1 | 0.2 | 0.3 | 0.4 | 0.6 | 0   | 0   | 0   | 0   | 0   | 0   |
|  | 1,000        | 0.1 | 0.2 | 0.4 | 0.6 | 0.8 | 1.0 | 0.1 | 0.1 | 0.2 | 0.3 | 0.5 | 0.7 | 0   | 0   | 0   | 0   | 0   | 0   |
| Well-sodded but overgrazed pastures and<br>meadows   | Entire slope | 0.1 | 0.2 | 0.3 | 0.4 | 0.6 | 0.7 | 0   | 0   | 0   | 0   | 0   | 0   | 0   | 0   | 0   | 0   | 0   | 0   |

1. Bare or fallow land free of vegetation will have erosion factors about double those given for intertilled crops. Idle land covered with weeds erodes about one-fourth to one-half as much as continuous intertilled crops.

2. The erosion factors for terraced land are based on experimental evidence which shows that properly constructed terraces with stable outlets can be expected to reduce erosion an average of about 80 per cent. On short gentle slopes, terraces will not reach this degree of effectiveness, whereas on longer slopes their effectiveness will approach 100 per cent. With upslope plowing the effectiveness of terracing is near 100 per cent for most rotations.

3. The erosion factors for contouring are also based on experimental evidence which shows that this practice, when considered for the entire year (on the average Missouri soil), can be expected to reduce erosion to at least 40 per cent less than that of the crop rotation alone.

4. Where slopes are over 12 per cent or longer than 1,000 feet, the factors should be increased in the same ratio as those listed for the more moderate or shorter slopes.

TABLE 45  
THE USE OF THE PRODUCTIVITY INDEXES AND EROSION FACTORS ON A 160-ACRE FARM \*

| Field No. or Letter | Acres in Field | Av. Slope (%) | Av. Length of Slope (feet) | Mechanical Erosion Control Measures | Crops Last Year                | Crops This Year            | This Year's Crop Use     | Kind   | Total used on † field | Soil Treatments |       | Productivity Index for Crops or Soil Treatments | Erosion Factors | Productivity Values P.I.† × Acres E.F.‡ |          |
|---------------------|----------------|---------------|----------------------------|-------------------------------------|--------------------------------|----------------------------|--------------------------|--------|-----------------------|-----------------|-------|---|-----------------|---|----------|
|                     |                |               |                            |                                     |                                |                            |                          |        |                       | (A)             | (B)   |   |                 | Positive                                | Negative |
| <b>A</b>            | 35             | 8             | 400                        | None                                | Permanent pasture              | Permanent pasture          | Grazed                   | Manure | 40 tons               | 0.00            | 0.40  | 6.00  | 14.00           |   |          |
| <b>B</b>            | 20             | 4             | 700                        | Terraced and contoured              | Wheat and 1st yr. sweet clover | 2nd yr. sweet clover, corn | Green manure All removed | None   |                       | +3.00           | -2.00 | 0.10  | 60.00           | 40.00                                   | 2.00     |
| <b>C</b>            | 20             | 6             | 500                        | Contoured                           | Wheat and lespedeza            | Oats and lespedeza         | All removed Hay          | None   |                       | -1.00           | 0.00  | 0.70  |                 | 20.00                                   | 14.00    |
| <b>D</b>            | 20             | 4             | 800                        | None                                | Oats and lespedeza             | Wheat and lespedeza        | Combined Grazed          | 4-16-4 | 4,000 lb.             | -0.65           | +1.30 | 1.20  | 26.00           | 13.00                                   | 3.60     |

|          |     |   |     |           |   |  |                          |                                |       |        |               |
|----------|-----|---|-----|-----------|---|--|--------------------------|--------------------------------|-------|--------|---------------|
| <i>E</i> | 20  | 6 | 600 | None      | Wheat and<br>1st yr.<br>timothy and<br>red clover | Hay<br>2nd yr. tim-<br>othy and<br>50% red<br>clover | Manure<br>50 tons        | 0.00<br>+0.15                  | 0.30  | 7.50   | 6.00          |
| <i>F</i> | 10  | 6 | 500 | None      | Alfalfa   | Hay  | 0-20-0                   | 2,000 lb.                      | +0.50 | 0.25   | 5.00<br>1.50  |
| <i>G</i> | 10  | 6 | 800 | Terraced  | Barley and<br>1st yr.<br>sweet clover             | Grazed   | None                     |                                | +2.25 | 0.00   | 22.50         |
| <i>H</i> | 20  | 4 | 400 | Contoured | Corn after<br>sweet clover                        | Wheat 1st<br>yr. sweet<br>clover                     | Combined<br>Left on land |                                | -0.65 |        | 13.00<br>8.00 |
|          |     |   |     |           |   |  | 0-20-0                   | 4,000 lb.<br>Manure<br>40 tons | +1.00 | 0.40   | 20.00         |
| <i>I</i> | 5   | 4 | 300 | None      | Garden, lots,<br>buildings                        | Garden, lots,<br>buildings                           | None                     |                                | 0.00  | 0.00   | 0.00          |
| Total    | 160 |   |     |           |   |  |                          |                                |       | 161.10 | 156.50        |

Productivity and erosion balance equals total positive of 161.1 and total negative of 156.5 or a positive of 4.6 for the whole 160 acres. Per cent gain or loss in productivity equals (total positive or negative, total acres) for this farm for the year,  $4.6/160$  or 0.03 per cent positive.

\* From Missouri *Exp. Sta. Bull.* 522.

† Each ton of manure is credited with a positive productivity factor of 0.15. Each 40 pounds of available plant food is credited with a positive productivity factor of 0.15.

‡ P.I. = productivity index; E.F. = erosion factor.

## 412 Fertility Maintenance and Productivity Rating

Frequently the sale price of land has been accepted as a measure of quality of land, notwithstanding that money value may be determined by factors other than crop productivity. However, there is a need for more exact comparisons or for some quantitative scale of ratings to be used in farm appraisal work, assessment of land for tax purposes, and in making inventories of land resources, which in turn are basic for soil conservation studies, rural zoning, and general land planning.

The desirability of having some scheme for the more exact comparison of the value or quality of soil and land, especially if constructed on a mathematical basis and expressed by indexes, being accepted, questions arise about the criteria to be used and their validity. Some of these questions are stated below.

### Questions

1. Should soils be rated on the basis of natural productivity alone, on the basis of management, or both?
2. How is a standard of value selected for a mathematical rating of productivity of soils?
3. What are the criteria which can be used in the classification of land on an agricultural basis?

**Basis for Rating Soils.** Soils can be rated solely on the basis of natural productivity or inherent fertility, but only by an assumption that management factors are constant. If the basis of management is emphasized or carried to the extreme, a rating of soils is likely to become nothing more than a comparison of the efficiency of different management practices, or a comparison of the relative efficiency of individuals or classes of men. Since the effect of the natural character of soil cannot be entirely eliminated by management practices, and since the productive capacity of soils may be greatly increased by the use of amendments, such as commercial fertilizers and by irrigation, it follows that both the natural productivity and the productivity obtained by amendments should be taken into consideration in making an index rating.

**Standard of Value.** Obviously, some standard of value has to be established in order to compare soil types on a mathematical basis. For example, the figure 100 may be taken as a standard, and this may represent the average yield for a particular crop in a region where the crop is extensively grown and where average yields are high. For example, the standard for corn may be 50 bushels per acre; then, if the average yields for a particular soil type are estimated at 25 bushels, the productivity rating for that soil type is 50. Soils may also be

given a "productivity grade." Productivity-grade numbers are obtained by a simple percentage weighting of the various crop indexes. Soils with a weighted average between 90 and 100 are rated 1; between 80 and 90, are rated 2; and so on. This is a general agricultural rating and not one for a particular crop. The method, here simply outlined, is in substance that employed by the Division of Soil Survey of the U. S. Department of Agriculture.

Table 46, showing the application of the method, is taken from a table of productivity ratings given in a Soil Survey report ("Soil Survey of Marshall County, Kentucky," United States Department of Agriculture, 1950). Contrasting soil types have been selected from the complete table for the county.

**Criteria for Rating Land.** *Land* should be distinguished from *soil* or *soil type* because these words, as generally used, are not exactly the same in meaning. If an individual soil type happens to be very uniform in character and occurs in large separate bodies, it can be taken as a satisfactory unit for land classification; however, in many places soils of widely divergent chemical and physical nature are closely associated in small bodies, and again configuration features of the landscape, especially the kinds of slopes, are an important factor affecting the use and value of land.

The possible criteria for the classification of land on an agricultural basis are (1) gross money value of agricultural products—plant and animal; (2) net income from land; (3) selling price of land; (4) values assessed for taxation; (5) measured yields of crops; (6) physical characteristics of the land.

Taken singly or used alone, none of the criteria listed above is entirely satisfactory except for some limited objective or for small areas of land. However, classifications, more or less subjective, including only a small number of divisions, 3 to 5, may be made and may have a considerable value in the inventory of land resources if a combination of factors, such as listed, is employed.

## Fertility Maintenance and Productivity Rating

TABLE 46

PRODUCTIVITY RATINGS OF SOME MARSHALL COUNTY, KENTUCKY, SOILS UNDER TWO LEVELS OF MANAGEMENT FOR CROPS  
COMMONLY GROWN

(Indexes in column *B* are the yields obtained under common management practices; those in column *C* are the yields that may be expected under good management practices.)

| Soils                             | Corn,<br>100 = 50<br>bushels |          |          | Wheat,<br>100 = 25<br>bushels |          |          | Lespedeza<br>Hay,<br>100 = 1 $\frac{1}{2}$<br>tons |          |          | Redtop<br>Hay,<br>100 = 1 $\frac{1}{2}$<br>tons |          |          | Soybean<br>Hay,<br>100 = 2 $\frac{1}{2}$<br>tons |          |          | Cotton<br>Lint,<br>100 = 400<br>lb. |          |          | Tobacco,<br>Dark,<br>100 = 1,000<br>lb. |              |              | Permanent<br>Pasture,<br>100 = 100<br>cow-acre<br>days |  |  | Principal Use |
|-----------------------------------|------------------------------|----------|----------|-------------------------------|----------|----------|--|----------|----------|---|----------|----------|--|----------|----------|-------------------------------------|----------|----------|---|--------------|--------------|--|--|--|---------------|
|                                   | <i>B</i>                     |          |          | <i>C</i>                      |          |          | <i>B</i>   |          |          | <i>C</i>  |          |          | <i>B</i>   |          |          | <i>C</i>                            |          |          | <i>B</i>                                |              |              |  |  |  |               |
|                                   | <i>B</i>                     | <i>C</i> | <i>B</i> | <i>C</i>                      | <i>B</i> | <i>C</i> | <i>B</i>   | <i>C</i> | <i>B</i> | <i>C</i>  | <i>B</i> | <i>C</i> | <i>B</i>   | <i>C</i> | <i>B</i> | <i>C</i>                            | <i>B</i> | <i>C</i> | <i>B</i>                                | <i>C</i>     | <i>B</i>     | <i>C</i>   |  |  |               |
| Huntington silt loam              | 80                           | 110      | *        | *                             | 85       | 120      | 75   | 85       | 80       | 90  | *        | *        | *  | *        | *        | *                                   | 65       | 65       | 90                                      | 90           | Corn, wheat, | Corn, wheat,   |  |  |               |
| Wheeling silt loam                | 60                           | 90       | 65       | 80                            | 65       | 120      | 55   | 80       | 55       | 80  | 90       | 130      | 80   | 95       | 80       | 55                                  | 55       | 80       | 80                                      | Corn, hay,   | hay          |  |  |  |               |
| Grenada silt loam,<br>level phase | 50                           | 75       | 45       | 55                            | 55       | 95       | 45   | 65       | 40       | 65  | 55       | 75       | 90   | 100      | 40       | 65                                  | 65       | 65       | 65                                      | Corn, hay,   | tobacco      |  |  |  |               |
| Sciotoville silt loam             | 50                           | 80       | 55       | 70                            | 60       | 100      | 45   | 75       | 50       | 70  | 80       | 110      | 80   | 95       | 45       | 70                                  | 70       | 70       | 70                                      | Corn, wheat, | hay          |  |  |  |               |
| Wheeling loamy fine<br>sand       | 50                           | 70       | 40       | 60                            | 40       | 75       | 40   | 60       | 40       | 60  | 75       | 100      | *  | *        | *        | 35                                  | 35       | 60       | 60                                      | Corn, hay    | Corn, hay    |  |  |  |               |
| Memphis silt loam                 | 40                           | 60       | 35       | 50                            | 40       | 75       | 25   | 45       | 30       | 50  | 45       | 70       | 60   | 70       | 30       | 30                                  | 30       | 30       | 30                                      | 30           | 30           | 30   |  |  |               |
| Henry silt loam                   | 25                           | 40       | *        | *                             | 40       | 65       | 35   | 60       | 30       | 50  | *        | *        | 65   | 80       | 25       | 25                                  | 25       | 25       | 25                                      | 25           | 25           | 25   |  |  |               |
| Lexington loam                    | 20                           | 40       | 10       | 35                            | 20       | 55       | 15   | 35       | 20       | 30  | 25       | 50       | 40   | 55       | 15       | 40                                  | 40       | 40       | 40                                      | 40           | 40           | 40   |  |  |               |

\* Crop not generally grown.

## Soils and Agriculture of Arid Regions

No generally accepted definition of arid region exists. A definition based solely on total annual precipitation is faulty because it does not take into consideration factors which influence the efficiency of a given amount of precipitation in the production of crops. Some of these factors are (1) distribution of the rainfall through the year, (2) temperature, (3) wind, (4) air humidity, and (5) amount of sunshine. A comparatively low rainfall, coming mostly during the growing season, may permit relatively high crop yields, while some areas, receiving as much as 70 to 80 inches of rain, with little of it coming during the growing season, require irrigation for crop production. Frequent fogs, moderate temperature, and little wind lead to low rates of evaporation of water from the soil and reduced transpiration of plants and hence reduce the moisture required for crop production.

For the purpose of this discussion the arid region is considered to include those areas with predominantly clear weather and annual precipitations as high as 15 inches, and also areas subject to fog, but with less precipitation, in which the soils have the same characteristics as those developed in the clear areas. Without irrigation, these lands are used principally for grazing, with a very small production of small grains in areas where conditions are especially favorable and under a system of one year of crop and one of fallow for moisture accumulation. Two headings under which arid soils may be studied are suggested.

### Objectives

- A. Characteristics and utilization of soil in arid regions.
- B. Development and management of saline and alkali soils.

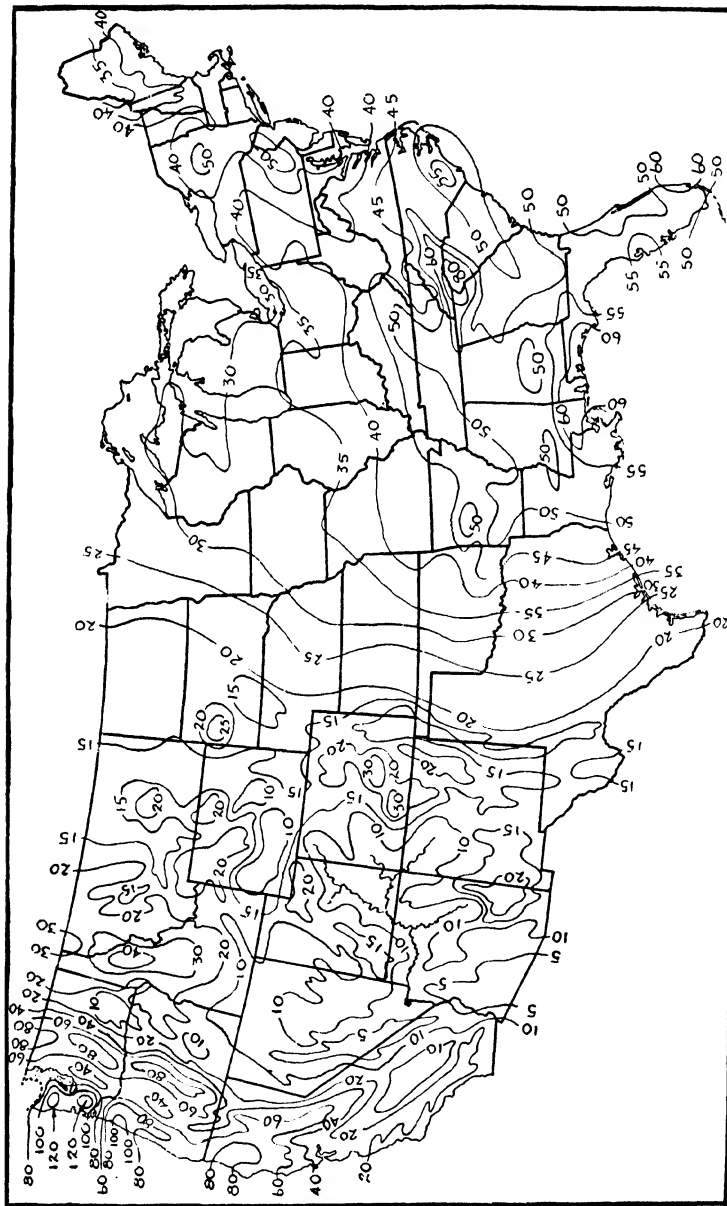


Fig. 88. A generalized map showing the total annual precipitation in different parts of the United States. Precipitation includes rain and the water equivalent of snow, sleet, and hail. Some of the smaller areas, such as those receiving 100 or 120 inches, are enlarged on the map. [Drawn from a map prepared by the Weather Bureau and presented in *Atlas of American Agriculture*.]

## CHARACTERISTICS AND UTILIZATION OF SOIL IN ARID REGIONS

Many misconceptions regarding soils of the arid regions have prevailed in the minds of laymen. These soils are not merely accumulations of more or less decomposed rock fragments with no profile characteristics, nor are they all of a coarse texture or sandy nature. They exhibit much diversity in properties. There are heavy soils, medium-textured soils, and sandy soils; young soils and soils with a pronounced profile development; alkali soils and those without injurious salt accumulations in the arid regions.

The agricultural utilization of arid soils is also varied. With an abundance of water for irrigation, intensive agriculture may be carried on, as described in the following chapter. Without irrigation, forage for grazing ranges from the very scanty supply of the true desert regions to the thin grass cover of the arid prairies. The following questions may serve as guides in studying these soils.

### Questions

1. What physiographic locations are occupied by arid soils?
2. What were the sources and nature of the parent materials for arid soils?
3. Which groups of soils are found in the arid regions?
4. What are the characteristics of red desert soils?
5. In what respects do gray desert soils differ from red desert soils?
6. How are sierozem soils described?
7. Are the deserts occupied by characteristic vegetation?
8. Does plant growth change desert soil?
9. Do the deserts make suitable grazing lands?
10. Are the deserts subject to erosion?
11. What are the characteristics of brown soils, and how are they utilized?
12. What are adobe soils?

**Locations of Arid Soils.** Arid soils of western United States occupy, in the main, broad valleys and basins between mountain ranges and extend up the slopes until increased precipitation or lower temperatures give rise to a different type of vegetation and to soils of different characteristics. In some places, however, arid conditions prevail over the entire surface of low mountain ranges, which are either bare of vegetation or are occupied by a thin cover of brush. The mountains serve as barriers to clouds and to cooling winds and so produce areas of low rainfall and relatively high mean annual temperatures. With

few exceptions, the relative air humidity is very low. Similar conditions prevail to a certain extent on the high plains lying directly east of the Rocky Mountains and give rise to areas classed in the brown soil group which are included among the arid soils in this discussion. In some locations reasons other than the occurrence of mountain barriers must be sought to explain the arid conditions, but a discussion

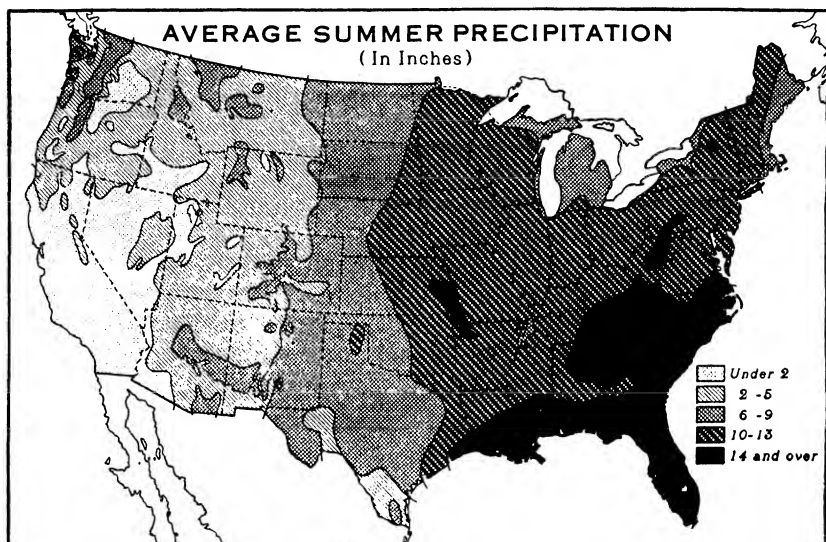


FIG. 89. Total annual precipitation does not always indicate whether or not crops can be grown without additional water. The amount of precipitation during the growing season is a better criterion. [From U.S.D.A. *Misc. Pub.* 260, by O. E. Baker. Drawn from a map prepared by the Weather Bureau and published in *Atlas of American Agriculture*.]

of these more unusual situations must be left to a textbook on climatology.

**Sources and Nature of Parent Material.** The basins in which the major portion of the arid soils has developed were occupied at some remote period by lakes or estuaries of the ocean. Evaporation of these waters has left deposits of salts or soil materials impregnated with salt. To a large extent these deposits have been covered with sediments carried by streams and ancient glaciers from the mountains. In fact, the true desert soils have developed mainly on long alluvial fans which stretch outward from the mountains until they meet the fans from adjacent parallel ranges. A belt of flat, low land, subject to

flooding and known as a playa, sometimes separates the fans. These alluvial fans provide the material for the development of deep, pervious, well-drained soils. The more recent alluvial deposits exhibit only the first stages of soil formation. Terraces of varying ages have furnished the material for soils with highly developed profiles. Some of the terrace soils have firmly cemented hardpans at varying depths, whereas dense claypans characterize the horizons of others.

The characteristics of these secondary soils are greatly influenced by the nature of the rocks furnishing the parent material. Granitic debris develops into pervious, medium-textured soils with desirable moisture relationships. Soils of heavier texture, which are sometimes less favorable for moisture infiltration, develop from the material contributed by basic igneous rocks. Sedimentary rocks may contribute to the formation of heavy clays, sandy soils, or soils of medium texture, depending on the proportions of shale, slate, sandstone, and limestone in the valley sediments.

The quantity of lime in the soil material has influenced the reaction of the horizons in the various profiles and has contributed to the formation of limestone nodules and of more or less cemented deposits of calcium carbonate known as *caliche*. The soluble salt content of the parent rocks and of the transporting streams has been influential in determining the alkalinity and the structure of the resulting soils.

More recent alluvium deposited by streams flowing through the lower portion of the valley floor may be rich in sand and silt, and highly pervious where it is dropped near the channel or fine-textured where water is impounded in depressions farther back.

On the foothills and mountain slopes residual soils of great variation in depth and texture are present. Their chemical and physical properties depend on the nature of the parent material and on moisture and temperature relationships.

In the following discussion no attempt is made to mention all the soil conditions occurring in the various areas considered. Attention is centered on the predominating soil-profile characteristics and on the accompanying vegetative cover and utilization practices.

**Arid Soil Groups.** Within the arid region, as defined in this chapter, are soils of the Gray and Red Deserts, Sierozems, and the Brown soils (except those approaching the dark brown or Chestnut soils) as described by Marbut.<sup>1</sup> The Desert soils usually occupy areas having less

<sup>1</sup> *Atlas of American Agriculture, Soils of the United States*, C. F. Marbut, U. S. Department of Agriculture, 1935, pp. 76-82.

than 7 to 8 inches of annual rainfall. The factor of the temperature resulting in different degrees of dehydration of hydrated iron oxides accounts for the division between the Gray and Red Deserts. Roberts<sup>2</sup> has observed that in Nevada the mean annual temperature of the Red Desert exceeds 60°F., whereas on the Gray Desert it is much less, possibly between 45° and 50°F.

The Brown soils occupy areas receiving 11 to 15 inches of rainfall, except in the fog belts of the Pacific Coast, where the precipitation may be as low as 10 inches.

**Red Desert Soils.** These soils are in the hot, arid regions covering the extensive southern part of Nevada, southeastern California, western Arizona south of the Grand Canyon, and sections of southern New Mexico and southwestern Texas. They vary in color from light pinkish gray to reddish brown and red. The upper subsoil in general is more compact and of heavier texture than the surface and is usually reddish brown or red. The lower subsoil is pink or white and very rich in lime. Maximum concentration of lime carbonate occurs at approximately 14 inches, although the zone may be many feet thick and frequently is consolidated. An intermixture of calcium sulfate often occurs below a depth of 2 feet. In many places the soil has blown or washed away sufficiently to leave a covering of stones on the surface, thus giving rise to the so-called "desert pavement." A coating of manganese and iron oxide on these stones gives them a polished appearance and hence is known as "desert varnish." Stones buried at shallow depths are coated on the underside with calcium carbonate, deposited by upward-moving water, and those at greater depths may be entirely covered with lime.

**Gray Desert Soils.** These soils lack the red or pink color found in the southern desert. They occupy the drier portion (less than 8 inches of rainfall) of the Great Basin, and areas in western Colorado and eastern Utah, with smaller areas in Idaho and Oregon.

The surface soil is calcareous and of a gray to light grayish brown color. The soil below, to a depth of several inches, is pale brown and grades into a more compact sometimes granular *B* horizon of heavier texture. The lime content increases with depth, forming a distinct zone at approximately 18 to 20 inches, which may take the form of a hardpan at this or greater depth. In both the Red and Gray Deserts the

<sup>2</sup> For this section much material has been taken from an unpublished manuscript by R. C. Roberts, Division of Soil Survey, Bureau of Plant Industry, Soils, and Agricultural Engineering, U. S. Department of Agriculture.

lime accumulations in the lower horizons may develop cemented formations known as caliche, which sometimes extend to a depth of many feet. Desert pavements develop where the soil is derived from material containing much coarse gravel and stone, but the desert varnish infrequently occurs.

**Sierozem Soils.** With increasing effective precipitation, the Gray Desert soils grade into the Sierozem soils, which comprise extensive areas in all eleven of the western states. They are confined to areas receiving approximately 8 to 11 inches of precipitation except in the north, especially along the Columbia River. There, owing to lower evaporation, the precipitation may be as low as 5 inches. The Sierozem soils have a light grayish brown or pale brown surface crust underlain by grayish brown or light brown laminated or vesicular layers 1 or 2 inches thick. This material abruptly changes to a brown or light brown granular, thin, heavier-textured layer which grades into a firm to compact brown layer. Free lime seldom occurs above a depth of 4 or 5 inches unless the parent material is highly calcareous. The layer of maximum lime accumulation occurs at a depth between 20 and 30 inches. The underlying material may be loose and friable, but many soils have lime hardpans comparable to that occurring in many of the Red and Gray Desert soils. The predominant shrub of the Sierozem soils is big sagebrush. Other vegetation includes juniper, black sagebrush and grasses such as blue grama, nigger-wool, galleta, and some crested wheat grass. The carrying capacity ranges from 8 to 20 head of cattle to the section.

**Vegetation of Red and Gray Deserts.** The characteristic vegetation of the Red Desert is creosote bush and bur-sage with relatively large numbers of many other plants, some of which are Wolfberry, Mormon tea, yucca, and prickly pear (in the northern part) and giant cactus, ocotillo, and ironwood (in the southern part). When the ground water table is nearer the surface, tamarack, mesquite, and cat's-claw occur. In areas with higher concentrations of salts are found iodine bush, quail bush, and salt grass.

On the Gray Desert shadscale, bud sage, rabbitbrush, and horsebrush are prominent. Among plants of importance for grazing purposes, white sage, winter fat, galleta, Indian rice grass, and bud sage should be mentioned. Alkali areas are characterized by the same plants found in similar locations, as on the Red Desert.

Vegetation on the deserts occupies only a small portion of the surface, seldom exceeding 20 per cent even in the most favored location and frequently almost disappearing. Possibly from 3 to 8 per cent coverage would be a reasonable average. It is noteworthy that grass

and other small succulent vegetation are found mostly around the base of the larger shrubs. This phenomenon is explained on the basis of the presence of a very thin crust underlain by a layer of soil containing innumerable small round pores filled with air (vesicular layer) which resists the entrance of water, except within the spread of the branches, and hence enough rain is absorbed under the shrub to supply



FIG. 90. In the desert, grass and other short succulent vegetation grow only around the base of larger shrubs.

the needs of both shrubs and smaller plants. Another explanation has been suggested by Brezeale, whose observations have led to the conclusion that the roots of the grass withdraw from the roots of the shrubs water which they in turn have obtained from the deeper subsoil.

**Soil Properties Modified by Desert Plants.<sup>3</sup>** Studies have shown that the soil under certain desert plants has a much higher *pH* value than soil in the areas between plants. Sometimes the increased *pH* is accompanied by a considerable increase in soluble salt content, and at times the increase in salts is small. In addition, there may be a decided increase in soluble salts without a great change in soil reaction. The greasewood appears to be more effective in increasing soil *pH* than other plants studied. Shadscale also raises the *pH* appreciably above that of bare soil. Tests show that the soluble salt content under pickleweed and samphire is quite high, whereas little salt accumulates under sagebrush and creosote bush.

<sup>3</sup> From a paper by Roy C. Roberts, prepared for the Fourth International Congress of Soil Scientists, July, 1950.

These phenomena are explained on the hypothesis that certain plants have greater power to absorb soluble salts or sodium from the various soil horizons than do others. The absorbed material is stored in the leaves and branches and is deposited on the soil as the leaves fall. The high *pH* values are due to the sodium so accumulated. Higher salt contents and greater *pH* values have been found under older rather than younger plants. This variation in soil reaction and salt content of soils under different species of plants and in bare areas must be taken into consideration when collecting soil samples for testing.

**Utilization of Desert Soils.** When supplied with water, many of the Desert soils are highly productive, as will be pointed out in the section dealing with irrigation. Without irrigation these are utilized almost entirely for grazing. The amount of vegetation produced which will be eaten by cattle, sheep, and horses is extremely meager, and hence carrying capacity on the year-round basis varies from virtually nothing to about 5 head of cattle per section on the Red Desert soils and possibly up to 10 head on the better-grassed northern portion of the Gray Desert. In general, the cattle are in better condition on the Gray Desert and graze in somewhat larger groups of about 10 or less. Sheep are more common on the Gray Desert, and as many as 1,000 may be herded together.

There is a strong tendency for Desert soils to form on the surface a thin, hard crust which retards the infiltration of rain. Under this crust a vesicular layer forms, sometimes to a thickness of 2 or 3 inches, which further reduces penetration of water. The development of these layers is more pronounced in the soils of heavier texture and, as previously mentioned, between the clumps of vegetation. As a result, the vegetative cover is more dense, especially for grass, on the soils of coarser texture, such as loamy sand and sands. These soils not only absorb a greater proportion of the rainfall but also give up a larger percentage of their moisture to plants. Furthermore, root penetration is deeper in such soils as the formation of hardpans of caliche is less frequent and, when present, is at a greater depth. In the non-irrigated areas deep, coarser-textured soils without a gravelly subsoil or substratum are accordingly the most highly prized on the desert.

**Erosion on the Desert.** Storms on the desert are frequently of torrential nature, which circumstance, coupled with the formation of crusts and layers of soil which resists infiltration of water, leads to severe erosion. Stream beds and dry washes, which through most of the year are entirely dry, carry torrents of water during storms and may wash

out highways or cover them with streams of such depth and velocity that they are impassable for many hours. Gully formation is common, and the washes are often of great depth. Sheet erosion removes the surface soil from the more gentle slopes. Water erosion is much more pronounced on the sandy loams and on soils of heavier texture than on the very sandy soils which are more receptive to rainfall.

The effects of wind erosion are evident everywhere. Each bush or clump of vegetation occupies a little mound, the soil of intervening spaces having been carried away to a greater or lesser extent. The formation of the desert pavement is an evidence of soil blowing as is the accumulation of wind-drifted sediment on the windward slopes of the hills and mountains. The giant sand dunes northwest of Yuma and extending some 50 to 60 miles in a northwesterly-southeasterly direction through southern California and northern Mexico are further proof of the erosive power of desert winds.

**The Brown Soil Area.** In the fog areas of the Pacific Coast, where about 80 per cent of the rainfall occurs from December to April, Brown soils develop with as low a precipitation as 10 inches. As the effect of fog and humidity diminishes, the soils are formed under higher precipitation until on the open plains and in the arid valleys they occupy areas receiving 11 to 15 inches. Although mean annual temperatures may reach 60°F. in the coastal zones, the soils, as a whole, are formed under temperatures of 45° to 50°F. Areas of Brown soil are found in both eastern and western Colorado, in western Nebraska, and in New Mexico, Wyoming, and Montana. Small areas are also scattered throughout the arid region where climatic conditions are somewhat too humid to permit of desert soil development.

The surface soil is grayish brown to brown and 6 to 8 inches in depth. The gray shade predominates over the brown of the humus to such an extent that sometimes it is difficult to distinguish the humus horizon from the underlying material. A thin, laminated layer has developed in the immediate surface. The *B* horizon is lighter in color than the surface soil (this is a distinguishing feature from the Desert soils), as is also the occurrence of cracks extending from the surface to the bottom of the *B* horizon. The horizon is friable and generally alkaline in reaction although not calcareous. Below the *B* horizon the soil is calcareous but is easily penetrated by roots and has a yellowish gray color with sometimes a brownish tinge. Maximum lime accumulation occurs at a depth of 2 or 3 feet, depending on moisture relations.

The dominant vegetation on the Brown soils is composed of various short grasses, of which blue grama, nigger-wool, bluegrass (*Poa* spp.),

June grass (*Koeleria cristata*), buffalo grass, and needle and thread grass are most prominent. Near the Pacific Coast wild oats and brome grass make their appearance. Several of the sages, prickly pear, rabbitbrush, and bitter brush, together with piñon pine and juniper on well-drained areas and cottonwood and willows near stream beds, are representative of the larger types of vegetation.

The vegetative cover is much more dense than on the Desert soils and affords a grazing capacity ranging from 20 to 35 cattle per square mile. Both cattle and sheep ranches are numerous. Stock raising is the main activity, although some dry farming is carried on. Water is usually supplied by means of windmills. Erosion is not so extensive as on the Desert soils, owing to the denser cover of vegetation, although much damage is evident on some of the more sloping fields that have been plowed for crop production.

**Adobe Soils.** The term adobe has been used with no great uniformity of meaning. The word is of Spanish or Spanish-American origin and originally referred to sun-dried bricks. More recently it has been applied to the soil from which the bricks were made. An examination by Hendry of bricks taken from old missions and other structures in southwestern United States revealed that the soils used were not of very fine texture, as often has been assumed, but varied from fine sandy loam to heavy loam. If the natural soil in the locality where the bricks were to be made was of too fine a texture, sand was added to make it more loam-like.

Replies to inquiries addressed to a considerable number of experiment stations and a study of soil survey maps and reports led Smith<sup>4</sup> to the following conclusions: Adobe soils occur primarily in the western states, having been mapped or described in Colorado, Wyoming, Idaho, Oregon, Utah, Nevada, California, Arizona, Texas, and New Mexico. They are derived from both transported and residual soil material, but not from wind-deposited sediment as has been sometimes suggested. The soils are developed in areas with an annual precipitation of less than 20 inches and in which there are distinct wet and dry seasons. Much adobe land is, and will continue to be, used for grazing, with possibly 5,000,000 acres being used for production of cultivated crops. Although a large variety of crops is grown, including citrus fruits, alfalfa, sugar beets, rice, and vegetables, the grain crops usually are grown to best advantage.

<sup>4</sup> "Characteristics of Adobe Soils," A. Smith, *Amer. Soil Survey Assoc. Bull.* 14, 1933, pp. 79-82.

The general conclusion of both Smith and Shaw is that adobe is a term which should be applied to heavy clay soils that exhibit a specific structural condition. Shaw<sup>5</sup> offered the following definition: "A soil which on drying cracks and breaks into irregular but roughly cubical blocks. The cracks are usually wide and deep and the blocks are from 20 to 50 or more centimeters across."

A comparison by Chang<sup>6</sup> of the physical properties of a number of adobe soils and of non-adobe clay soils of similar clay content led to the following conclusion. Upon drying, adobe soils shrink extensively, developing wide, straight cracks which divide the soil into relatively large blocks of primary structural units. These blocks as well as the smaller secondary units into which they break have approximately flat sides, few in number, which meet at angles approaching right angles. In other words, the structural units are roughly cubical. They were found to have a high resistance to crushing. Soils containing a large quantity of montmorillonitic clay in a dispersed condition have an adobe structure unless other factors interfere. On the other hand, clay soils containing kaolinitic clay or montmorillonitic clay in a flocculated condition are unlikely to develop such a structure. Some factors which tend to prevent the formation of adobe structure are the presence of a considerable quantity of (1) soluble salt, (2) calcium carbonate, (3) organic matter, and (4) sand.

#### DEVELOPMENT AND MANAGEMENT OF SALINE AND ALKALI SOILS

Saline and alkali soils may be defined as soils containing sufficient soluble salts to impair their productivity. The quantity, proportion, and nature of the salts present may vary in the two groups. The soils under discussion may be classified in three groups, namely saline soils, saline-alkali soils, and non-saline-alkali soils.<sup>7</sup> Several questions will serve as guides in the study of these soils.

<sup>5</sup> "A Definition of Terms Used in Soil Literature," C. F. Shaw, in *Proceedings and Papers of the First International Congress of Soil Science*, Com. V, Vol. 4, 1927, p. 57.

<sup>6</sup> "An Experimental Study on the Development of Adobe Structures in Soils," C. W. Chang, *Soil Science*, Vol. 52, 1941, pp. 213-227.

<sup>7</sup> This classification and description of the soils was derived from *Diagnosis and Improvement of Saline and Alkali Soils*, U. S. Regional Salinity Laboratory, Riverside, California.

## Questions

1. What is meant by saline soils?
2. How do saline-alkali soils differ from saline soils?
3. Why are alkali soils so designated?
4. What conditions give rise to saline and alkali soils?
5. Where do the salts come from that cause salinity and alkalinity?
6. Why are high concentrations of soluble salts detrimental to many plants?
7. How does alkalinity affect soil conditions?
8. Which plants are most tolerant of alkalinity and salinity?
9. How may saline soils be reclaimed?
10. Are special treatments needed for the reclamation of "black alkali" soils?
11. How does removal of alkali affect soil structure?

**Saline Soils.** These soils contain a relatively high concentration of soluble salts, made up largely of chlorides, sulfates, and sometimes nitrates. Small quantities of bicarbonates may occur, but soluble carbonates are usually absent. Frequently relatively insoluble salts, such as calcium sulfate, and calcium and magnesium carbonates are also present. The chief cations present are calcium, magnesium and sodium, but sodium seldom makes up more than one-half of the soluble cations and is not adsorbed to an appreciable extent on the colloidal fraction of the soil.

The pH value of these soils is 8.5 or less, and the exchangeable sodium percentage is less than 15. White crusts frequently accumulate on the soil surface, and streaks of salt are sometimes found within the soil. Saline soils have a favorable structure because the colloids are highly flocculated. Soils in this group are similar to those designated as "white alkali" soils in older publications.

**Saline-Alkali Soils.** Soils in this group are also characterized by a high concentration of soluble salts, but they differ from saline soils in that the exchange sodium percentage is greater than 15. As long as the large quantity of soluble salts remains in the soil, the high sodium content in the colloids causes no trouble and the soil pH seldom exceeds 8.5. If, however, the soluble salts are temporarily leached downward, the pH goes above 8.5, the sodium causes the colloids to disperse, and a structure unfavorable for tillage, the entry of water, and root development develops. The movement of the soluble salts upward into the surface soil may lower the pH and restore the colloids to a flocculated condition. The management of this group of soils is a problem until the excess soluble salts and the exchangeable sodium are removed from

the zone of root growth. Unless calcium sulfate or another source of soluble calcium is present, the drainage and leaching of these soils convert them into non-saline-alkali soils.

**Non-Saline-Alkali Soils.** These soils do not have so high a concentration of soluble salts as do those in the two preceding groups, but they have a larger amount of adsorbed sodium; the exchangeable sodium percentage exceeds 15. The *pH* values usually range between 8.5 and 10. The sodium hydrolyzes from the colloids, and small amounts of sodium carbonate may form. Organic matter in the soil is highly dispersed and is distributed over the surface of the particles, giving a dark color; hence the term "black alkali" which was formerly used to designate such soils. These soils frequently occur in small irregular areas in regions of low rainfall and are referred to as "slick spots."

Alkali soils commonly develop as a result of irrigation. Because of the dispersed state of the colloids, the soils are difficult to till and are slowly permeable to water. After a long period of time the dispersed clay may migrate downward, forming a very dense layer with a prismatic or columnar structure. When this phenomenon occurs, a few inches of relatively coarse-textured soil may be left on the surface.

The soil solution of non-saline-alkali soils contains only small amounts of calcium and magnesium but larger quantities of sodium. The anions include sulfate, chloride, bicarbonate, and usually small quantities of normal carbonate. In some areas an appreciable amount of potassium salts is also present.

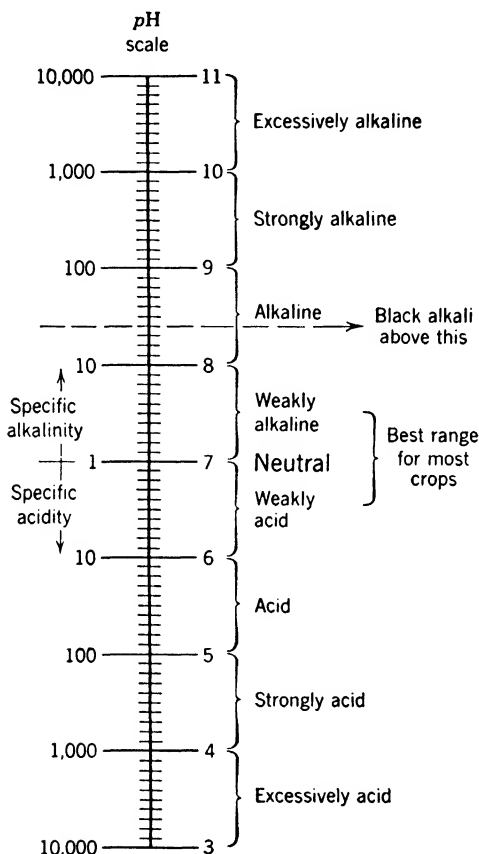
**Formation of Saline and Alkali Soils.<sup>8</sup>** The ground water of arid regions usually contains considerable quantities of soluble salts. If the water table is high, large amounts of water move to the surface by capillarity and are evaporated, leaving an ever-increasing accumulation of soluble salts. Through this process the upper parts of the soil may become so highly impregnated with salts that only alkali-resistant plants will grow. In locations where the water table is too deep for appreciable capillary movement to the surface, harmful accumulations of soluble salts will not develop. The rate of salt accumulation will be determined by (1) the rate of capillary movement of water to the surface, (2) the salt content of the ground water, and (3) the rapidity of evaporation. The nearer the water table is to the surface, the more rapid will be the salt accumulation, other factors being the same.

Under irrigation, alkali soils have been developed by several means.

<sup>8</sup> Many of the ideas in this section were obtained from "The Reclamation of Alkali Soils," W. P. Kelley, *California Agr. Exp. Sta. Bull.* 617, 1937.

(1) Excessive applications of water have raised the ground water level sufficiently to permit concentration of salts through evaporation. (2) Seepage from leaky canals and lateral ditches has resulted in a high

**Chart 9. A Chart Showing the Relationship between Soil Reaction (*pH*) and Alkalinity**



[Supplied by W. T. McGeorge]

water table. (3) The use of irrigation water with a high salt content has resulted in salt accumulation when (a) drainage is poor so that the salts cannot be leached out, (b) the application of water is so limited that the salts are left in the root zone in place of being washed out, and (c) the water contains such quantities of sodium salts that the colloids become impregnated with them and the favorable structure of the soil is destroyed.

Alkali spots may also develop in irrigated fields when for some reason a given area of soil does not absorb the water. Water that is carrying salts will move into such soil from the sides and below, and through surface evaporation salts will accumulate. Because no water enters from the surface, the situation becomes steadily worse and the spots may enlarge.

The overflow of streams by flood waters may cause a rise in the ground water level of adjacent valleys or may also cause the seepage of flood water through the porous stream bed. Again the irrigation of near-by land of higher level may result in a higher water table. All such increases in height of ground water carry the possibility of alkali development.

**Sources of Alkali Salts.** Alkali salts are derived primarily from the soil and rocks through which percolate the waters that supply the water table and the irrigation streams. The salt supply of the rocks and soil may be the result of accumulation when the area was a portion of the ocean bed or of the bottom of a saline lake, as in sedimentary deposits. The chemical decomposition of minerals has also contributed some salts. In igneous rocks, chemical weathering has supplied the soluble material. Kelley has observed that alkali soils, watered with drainage from sedimentary rocks, contain a higher proportion of sulfates than do those receiving water from igneous rocks. On the other hand, more black alkali soils appear to develop when the water supply comes from an area of igneous rocks. The primary source of the sodium, calcium, magnesium, and sulfur is evident, for many minerals contain these elements. The source of the sodium carbonate is somewhat more difficult to explain in regions of sparse vegetation where the production of carbon dioxide from decay of organic matter and respiration of plant cells is limited. Some sodium carbonate may be produced through the interaction of sodium chloride or sodium sulfate with calcium carbonate. It should be remembered, however, that the quantity of sodium carbonate present even in black alkali soils is not great.

**Detrimental Effects of Alkali on Plants.** High concentrations of neutral salts such as sodium chloride and sodium sulfate may interfere with the absorption of water by plants through the development of a higher osmotic pressure in the soil solution than exists in the root cells. Furthermore, the wilting coefficient of soils is raised by salt accumulations, and hence the quantity of water a soil will supply to plants may be reduced through the presence of alkali. Detriment to plants may result also from soluble salts when the concentration is not sufficient to influence absorption of water. The entrance of nutrient ions into root

hairs is influenced by the nature and concentration of other ions present. Alkali salts may therefore result in nutritional difficulties in crops because of their inability to absorb needed nutrients from the soil.

Black alkali is much more harmful to plants than is white alkali. The highly alkaline reaction due to the presence of sodium carbonate and the large quantity of adsorbed sodium represses the availability of several nutrients, especially iron, manganese, and phosphorus. Also, the alkaline soil solution has a corrosive action on the bark of roots and stems. The high concentration of neutral salts, which usually occurs in black alkali soils, affects the absorption of moisture by the plants just as it does in white alkali soils.

**Detrimental Effects of Alkali on Soils.** As previously mentioned, the sodium in black alkali soils results in a deflocculation of the colloids and hence in a breaking down of the soil structural units. This puddled condition renders the soil more or less impervious and retards entrance of irrigation and rain water and impedes drainage. In heavy-textured soils the penetration of roots may be restricted by the density of the deflocculated zone. Aeration is also much reduced, setting up anaerobic conditions and resulting in the formation of reduced compounds which are toxic to plants.

The presence of certain white alkali salts, largely those of calcium, tends to counteract to some extent the detrimental effects of the adsorbed sodium and the sodium carbonate. The general action of the neutral salts is to flocculate the colloids and hence preserve the normal soil structure.

**Tolerance of Plants for Alkali.** Plants differ markedly in their tolerance of alkali, and likewise the alkali salts differ greatly in their detrimental effects on plants. Hilgard set the limit of tolerance for the principal salts found in alkali soils as follows:

Sodium carbonate, 0.10 to 0.25 per cent

Sodium chloride, 0.20 to 0.50 per cent

Sodium sulfate, 0.5 to 1.00 per cent

Unquestionably, associated conditions, such as temperature, moisture supply, organic matter content, soil texture, and supply of nutrients, influence the tolerance of plants for these salts.

Shantz<sup>9</sup> lists some of the most alkali-resistant native plants, and the percentage of salts in soils in which they are found growing, as follows:

<sup>9</sup> "Plants as Soil Indicators," H. L. Shantz, *U.S.D.A. Yearbook* 1938, p. 852.

- Greasewood, more than 0.5 per cent
- Seepweed, 2.5 per cent or more
- Pickleweed, 1.0 to 1.5 per cent
- Salt grass, somewhat less than 1.0 per cent
- Alkali sacaton, about 0.5 per cent

The creosote bush of the southern deserts and sagebrush of the northern deserts indicate soils containing harmless quantities of salts. Dense stands of desert salt bush occur on fine-textured soils containing some salt, but an amount insufficient to damage crops under irrigation unless further salt accumulation is permitted.

The relative tolerance of a number of fruits and field and truck and forage crops to soluble salts is indicated in Chart 10. The crops listed first in each group are considered more tolerant, and those named last most sensitive, to salinity.

#### Chart 10. The Relative Tolerance of Several Crops for Salinity \*

| Good Salt Tolerance          | Moderate Salt Tolerance     | Poor Salt Tolerance      |                    |
|------------------------------|-----------------------------|--------------------------|--------------------|
| <i>Fruit Crops</i>           |                             |                          |                    |
| Date palm                    | Pomegranate                 | Pear                     | Apricot            |
|                              | Fig                         | Apple                    | Peach              |
|                              | Olive                       | Grapefruit               | Orange             |
|                              | Grape                       | Prune                    | Lemon              |
|                              |                             |                          | Almond             |
|                              |                             |                          | Avocado            |
| <i>Field and Truck Crops</i> |                             |                          |                    |
| Barley (grain)               | Rye (grain)                 | Broccoli                 | Radish             |
| Sugar beets                  | Wheat (grain)               | Tomato                   | Celery             |
| Milo                         | Oats (grain)                | Cabbage                  | Beans              |
| Rape                         | Rice                        | Cauliflower              |                    |
| Cotton                       | Alfalfa                     | Lettuce                  |                    |
| Beets (garden)               | Sorghum (grain)             | Sweet corn               |                    |
| Kale                         | Corn                        | Carrot                   |                    |
| Asparagus                    | Foxtail Millet              | Onion                    |                    |
| <i>Forage Crops</i>          |                             |                          |                    |
| Alkali sacaton               | White sweet clover          | Wheat (hay)              | White Dutch clover |
| Salt grass                   | Yellow sweet clover         | Oats (hay)               | Meadow foxtail     |
| Nuttall alkali grass         | Perennial rye grass         | Orchard grass †          | Alsite clover      |
| Bermuda grass                | Mountain brome              | Blue grama               | Red clover         |
| Rhodes grass                 | Barley (hay)                | Meadow fescue            | Ladino clover      |
| Rescue grass                 | Birdsfoot trefoil           | Reed canary              | Burnet             |
| Canada wild rye              | Strawberry clover           | Big trefoil              |                    |
| Beardless wild rye           | Dallis grass                | Smooth brome             |                    |
| Western wheat grass          | Sudan grass                 | Tall meadow oat<br>grass |                    |
|                              | Huban clover                | Cicer milk vetch         |                    |
|                              | Alfalfa (California common) | Sour clover              |                    |
|                              | Tall fescue                 | Sickle milk vetch        |                    |
|                              | Rye (hay)                   |                          |                    |

\* From "Diagnosis and Improvement of Saline and Alkaline Soils," U. S. Regional Salinity Laboratory, Riverside, California, 1947.

† Salt tolerance of remaining crops is "fair" rather than "moderate."

**Reclamation of Saline or White Alkali Soils.** Good drainage is necessary for the reclamation of alkali soils. It is essential in the reclamation process to remove the excess salts from the root zone, and this can only be done by the application of sufficient water to wash them into the lower soil depths. Unless there is ample drainage, the addition of so much water will raise the water table and hence lead to increased accumulations of salt in the surface soil rather than to a correction of the alkali condition. Sufficient drainage should be provided to reduce the ground water level well below the zone of root penetration. Kelley states that "preferably the ground water should never be less than 8 to 10 feet below the surface and every reasonable effort should be made to prevent its rising nearer than 5 to 6 feet from the surface even for brief periods."

With ample drainage provided, one may proceed to the leaching out of the salts. In fine-textured soils the reclamation process will be slow, and doubly so if the soil is underlain by a heavy clay subsoil. In fact, the presence of a dense clay layer makes difficult the removal of salts from even medium- or coarse-textured soils. It is questionable if reclamation of soils with very deep, heavy clay subsoils is feasible from an economic standpoint.

Experiments have shown that leaching is all that is needed to reclaim white alkali soils that have sufficient drainage. The addition of chemicals or plowing under of manure or green-manuring crops is unnecessary. No specific directions can be given regarding the frequency of irrigation or the quantity of water to apply at each irrigation. The main points to observe are (1) that the soil be kept moist so that the soil solution will not become sufficiently concentrated to damage the growing crop, (2) that sufficient water be applied at each irrigation to result in some leaching of salts into the drainage water, and (3) that the soil of each irrigation check be carefully leveled so that the water will enter the soil uniformly.

**Reclamation of Black Alkali Soils.** All that has been said concerning the need for drainage and the application of sufficient irrigation water to cause leaching is of as much, if not more, importance in the reclamation of black alkali soils than in the treatment of white alkali. Although it has been shown that the application of ample irrigation water, coupled with good farming practices, will ultimately result in the removal of black as well as white alkali, the reclamation process may be materially hastened through the application of various chemicals. The basis of the treatments is the replacement of adsorbed sodium in the colloidal fraction by calcium and the conversion of the

replaced sodium and any occurring as the carbonate into neutral sodium sulfate.

The desired changes may be brought about by applications of considerable quantities of finely ground calcium sulfate (gypsum). Ground sulfur, however, will accomplish the same results somewhat more quickly, and the application need not be heavy, 1,000 to 2,000 pounds per acre having sufficed for some soils. The sulfur is oxidized in the soil to sulfur trioxide, which combines with water to make sulfuric acid. Other soluble sulfates such as iron or aluminum have also proved effective. As a supply of soluble calcium is needed to complete the reactions, additions of manure or the plowing under of a green-manuring crop, in addition to the chemical treatments, are helpful. The decaying organic matter produces CO<sub>2</sub>, which, with H<sub>2</sub>O, combines with CaCO<sub>3</sub>, usually present in alkali soils, to form the more soluble Ca(HCO<sub>3</sub>)<sub>2</sub>.

The growth and plowing under of sweet clover or the applying of manure is a good way to start the reclamation process whether or not chemicals are to be applied. The growing of Bermuda grass, a highly alkali-resistant plant, also has proved an advantageous method of starting alkali removal practices. The data in Table 47 show the effects of

TABLE 47

EFFECTS OF VARIOUS SOIL TREATMENTS ON THE YIELDS OF CROPS GROWN ON  
BLACK ALKALI SOIL.\*

(Yields in pounds per acre)

| Date   | Crop    | Sulfur            |                   |                   |                     | Sulfur,<br>2 Tons |
|--|---------|-------------------|-------------------|-------------------|---------------------|-------------------|
|  |         | Sulfur<br>2 Tons, | 2 Tons,<br>Ground | Gypsum<br>2½ Tons | Limestone<br>2 Tons |                   |
| (Sweet clover plowed under as green manure on all plots) |         |                   |                   |                   |                     |                   |
| 1926   | Alfalfa | 10,554            |                   | 11,129            | 80                  | 11,129            |
| 1927   | "       | 16,984            |                   | 18,291            | 596                 | 18,243            |
| 1928   | "       | 18,953            |                   | 20,323            | 1,693               | 19,679            |
| 1929   | "       | 10,484            |                   | 14,113            | 5,161               | 15,080            |
| 1930   | "       | 5,000             |                   | 6,450             | 2,095               | 5,565             |
| 1931   | "       | 12,661            |                   | 15,242            | 9,596               | 14,233            |
| 1932   | "       | 14,033            |                   | 15,161            | 14,516              | 18,106            |
| 1933   | "       | 12,422            |                   | 16,814            | 16,885              | 18,549            |
| 1934   | "       | 12,097            |                   | 16,289            | 17,823              | 18,385            |
| 1935   | Oat hay | 2,742             |                   | 3,790             | 3,838               | 4,597             |
| 1936   | " "     | 6,129             |                   | 5,645             | 6,613               | 6,853             |

\* "The Reclamation of Alkali Soils," W. P. Kelley, *California Agr. Exp. Sta. Bull* 617, 1937, p. 13.

various treatments in the reclamation of a black alkali soil in California, as measured by the yields of crops. The treatments were applied in May, 1923, after which the soil was allowed to lie idle, with an occasional light irrigation and cultivation, until February, 1925. After the plot was flooded twice to leach out salts, white sweet clover was seeded and the crop plowed under as green manure in September. The next February alfalfa was seeded.

It will be noted that, although the yields on the untreated plot were quite low at first, they were very satisfactory after 6 years. So successful have been experiments in the reclamation of alkali soils that both Kelley of California and McGeorge of Arizona, the two men who have probably done more work on this problem than any other living investigators, maintain that virtually any such soil may be reclaimed if adequate drainage can be established and if the soil is not underlain by a very difficultly pervious clay subsoil. They are of the opinion, furthermore, that any soil under irrigation may be maintained sufficiently free of alkali and in a satisfactory structural condition if recommended management practices are followed, particularly those concerning irrigation.

**Effect of Alkali Removal on Soil Structure.** Black alkali soils are of notoriously poor structure, becoming very hard upon drying and breaking up into clods, which are very difficult to crush, when plowed. When a considerable quantity of white alkali salts accompanies the black alkali, the structural condition is not so bad, and a satisfactory seedbed may be prepared more easily. The leaching out of the alkali tends to leave the soil in an even worse physical condition, and hence some investigators have advised against this method of reclaiming such soils. Experiments have shown, however, that applications of calcium, iron, or aluminum sulfates, and also sulfur, followed by proper irrigation, tend to flocculate the colloids and cause the rapid development of a granular structure. Soils from which the alkali is leached without the application of any of the chemicals mentioned are much slower in developing a granular structure. The growing of a crop such as Bermuda grass or the plowing under of a green-manuring crop hastens the granulation process.

Although the structure of white alkali soils is impaired by leaching, the detriment is not so great as in black alkali. The growth of Bermuda grass or of alfalfa for a few years or the plowing under of green-manuring crops rapidly improves the soil structure.

# 19

## Irrigation

The presence of remnants of canals in the Southwestern States indicates that irrigation was practiced there by the prehistoric peoples who once inhabited that region. So accurately were some of these canals located with respect to grade and convenience in water use that ditches of modern irrigating systems either follow or parallel them.

According to a special release, the 1945 Census shows 20,539,470 acres of irrigated land in farms in 1944. This is an increase of more than two and one-half million acres over the 1939 acreage. This increase has resulted in part from the development of new irrigation projects, both private and public. Bureau of Reclamation projects in Arizona, Colorado, Idaho, Montana, and Nevada have added approximately 170,000 acres. There are some nineteen states in which irrigation is of relatively great importance. The number of farms reporting irrigated land dropped from 299,604 in 1939 to 288,195 in 1944. These figures indicate some consolidation of farms into large units.

There has also been a marked increase in acreage of irrigated land in the semi-arid and humid portions of the country. For example, the 1945 Census reports the acreage of irrigated land in 1939 and 1944, respectively, to be: in New England, 2,848 and 12,155; in the Middle Atlantic States, 17,260 and 30,792; in the east North Central States, 10,833 and 12,795; and in the South Atlantic States, 128,037 and 224,446. Many experiments are under way to determine the feasibility of irrigating various field crops and pasture in different sections of the non-arid parts of the country. The subject of irrigation will be considered under two general headings.

### Objectives

- A. Water supply and land for irrigation.
- B. Irrigation practice.

**WATER SUPPLY AND LAND FOR IRRIGATION**

Irrigation diminishes one of the greatest hazards in crop production, namely, inadequate water supply. In few instances is this factor entirely eliminated, however, as a shortage of water during some part of the growing season is not an uncommon occurrence on many irrigation projects. Also, breaks in canals as a result of floods and destruction of water control structures by fire may require the turning off of the water with resultant crop damage. The right to obtain water from streams and canals for irrigation purposes and the cost of the water are points to be considered. Furthermore, all land is not well adapted to irrigation and considerable cost for preparation may be involved. Answers to the following questions are pertinent in selecting land for farming under irrigation.

**Questions**

1. Do farmers have a right to take water from streams crossing their lands?
2. How does one procure a water right?
3. What precautions should be taken in buying land for irrigation and in bargaining for water?
4. What kind of land is suitable for irrigation?
5. Is the preparation of land for irrigation costly?
6. Why is drainage essential in irrigation?
7. What points should be considered in constructing irrigation ditches?

**Obtaining Water from Streams and Canals.** Water from streams is usually appropriated to individuals and canal companies according to priority rights, determined by the date the arrangements are made for water and the supply available. When two or more companies or individuals obtain water from a stream which is subject to wide variations in flow, the last to obtain water rights is the first to suffer when a scarcity arises. Although in some states the doctrine of riparian rights is recognized—that is, the owning of land along a stream entitles one to withdraw water from the stream for irrigation purposes—this right is modified by the doctrine of appropriation. In most of the western states the appropriation, diversion, and distribution of water from streams are under the direction of a state engineer or irrigation board.

**Procurement of Water Rights.** The present-day settler or purchaser of irrigated land usually bargains with a canal company or water users' association for his water supply. In the early days of irrigation

a perpetual water right was purchased for each tract of land, and an additional annual charge was made for operation and maintenance of the canal system. At present the more common procedure is the sale of rights, along with the land, which carry an interest in the water-supplying system. Under this plan the system becomes the property of the landowners when a certain proportion of the rights has been paid for. Another method is the organization of an irrigation district which includes all land to be watered from a given system. Under this arrangement no water right is purchased, as ownership of land carries the right to water. The cost of the irrigation system, however, is levied against the land in the form of taxes.

When the government develops an irrigation project and supplies water to privately owned land, it operates through a water users' association. All the landowners become shareholders in the association through the purchase of water rights, the cost of which varies from \$25 to \$90 per acre. This cost is payable in 20 annual instalments. The operation and management of the irrigation system, exclusive of storage reservoirs, usually passes into the hands of the landowners when payment of water rights for the major portion of the land has been made. The maintenance and operation of the system then becomes the responsibility of the landowners, and the cost is prorated among them.

**Precautions in Buying Land and in Bargaining for Water.** Although it is assumed that no private company or water users' association will contract to irrigate more land than the supply of water is adequate for, it is well to investigate the following points before closing a deal. (1) What "right" does the land have to water? Was it among the first or the last to obtain rights to water from the present source? (2) What right, or priority of right, to water has the company with which you are bargaining? (3) Is the source of water adequate to supply crop needs during critical periods? Sometimes there is an abundance of water during flood seasons but a very limited supply at other times. Is there water for the last irrigation of late-growing crops like alfalfa or potatoes? (4) Is the cost of water stabilized, or may there be a decided increase in cost in years to come?

**Selecting Land for Irrigation.** In choosing land for irrigation, a careful examination should be made of the soil to determine (1) texture of soil to depth of several feet; (2) presence of a difficultly pervious stratum or of gravel within a depth of 5 or 6 feet; (3) accumulation of soluble salts in injurious quantities; (4) slope and evenness of soil surface; and (5) behavior of the soil under irrigation. A desirable soil is readily pervious to water and yet is moisture-retentive. It is well

if the soil will absorb sufficient moisture in 24 hours to wet it to a depth of 2 or 3 feet. Some soils are so slowly pervious that they become wet to a depth of only a foot or less in 24 hours. On the other hand, some soils are so coarse-textured that water passes rapidly below the reach of plant roots, and little available moisture is retained.

A difficultly pervious stratum within the root zone will limit root penetration and may result in the waterlogging of the overlying soil. On the contrary, a stratum of gravel or coarse sand in the root zone may result in the loss of considerable irrigation water and in the storage of a limited supply for crop use.

Usually it is possible to judge the behavior of the soil under irrigation by observing similar soil in near-by irrigated fields. If this cannot be done, water may be applied to a small area for a trial. In general, if the soil is too heavy, it will absorb water slowly, be sticky and hard to cultivate when damp, and will crack and bake on drying, although some very heavy soils, particularly those with an adobe-like structure, absorb water readily and are not difficult to manage. Some sandy soils, on the other hand, form plow soles<sup>1</sup> very readily. Soil of too coarse a texture will absorb water very rapidly and show little coherence when dry.

In the absence of chemical tests, some knowledge of the presence or absence of alkali salts may be obtained by observing the natural vegetation. This point is discussed in the section dealing with alkali soils.

The land surface should be comparatively smooth, for the cost of leveling land is high. A uniform slope of 10 to 20 feet to the mile is desirable, although much steeper slopes are in use. Land cut up by ravines, gullies, or buffalo wallows, or covered with sand dunes and hummocks should be avoided if possible.

**Cost of Land Preparation for Irrigation.** The cost of preparing land for irrigation varies widely. If the surface is very smooth and the slope uniform and of desirable grade, the cost is very little. On the other hand, if there is much leveling to be done, boulders to be removed, ditches to be built over rough topography, and similar improvements to be made, the cost per acre may be quite high.

**Providing Drainage.** It is essential to have good drainage if irrigated land is to remain permanently productive. If the land is not naturally well drained, drainage ditches must be provided before the

<sup>1</sup> A plow sole is a layer of soil in which the clay has become deflocculated and the soil packed into a dense layer as a result of repeated plowing to the same depth.

land is in use many years. It is difficult to apply sufficient irrigation water for maximum crop growth without getting some excess. In fact, some excess water frequently is desirable in order to wash out alkali salts. If the drainage water is not removed, it may raise the water level, thus increasing the danger of alkali accumulation. Furthermore, the subsoil may become waterlogged with a resultant restriction of root development and crop damage.

**Irrigation Ditches.** The main canals of an irrigation system are constructed and maintained by the company or association. The ditches carrying the water to and over individual farms, however, are constructed and maintained by the farmers themselves.

The capacity, location, and design of farm ditches depend on several factors, among which the following deserve especial consideration: (1) depth and permeability of the soil; (2) whether the water is delivered in a continuous stream or for short periods with long intervals between; (3) the method to be used in applying the water to the land; (4) the acreage to be irrigated; and (5) the water requirements of the crops to be grown.

If the soil is very permeable, there is much loss of water by seepage unless it contains sufficient silt and clay to form a seal on the sides and bottom of the ditch. In extreme cases pipes are used in place of ditches. These pipes usually are of concrete or of wood staves held together by means of bands applied by machinery. If redwood is not used, the staves should be treated with preservatives to delay decay.

If water flows continuously, smaller ditches will suffice than if all the water needed must be obtained during a limited time at infrequent intervals. Likewise, large acreages and crops which require much water necessitate the construction of large ditches. In general, ditch capacities range from less than 1 to more than 10 cubic feet per second. Methods used to supply water to the land will be discussed in another section.

Water is carried across ravines, gullies, or depressions by one of three methods: (1) wood or metal flumes are built on grade across the obstruction; (2) a pipe may be laid in the depression in the form of an inverted siphon; (3) if the depression is only a few feet deep, levees may be built on each side.

#### IRRIGATION PRACTICE

Artificial watering adds certain complexities to farming practice. Judgment and experience are necessary to know when to apply water

and how much to apply for different crops. Irrigation offers the opportunity for distribution of weed seed over the fields. Likewise, improper watering and irrigation without adequate drainage may result in alkali accumulation. Water costs and the maintenance of ditches are also important considerations in the expense of crop production.



FIG. 91. A "well-built wooden flume." [From U.S.D.A. *Farmers' Bull.* 1348, by James C. Marr.]

Several questions are offered to serve as guides in studying the problems involved in irrigation farming.

### Questions

1. How may irrigation water be applied?
2. What quantity of water is needed for adequate irrigation?
3. How can a suitable irrigation system be selected?
4. By what systems is irrigation water measured?
5. What provisions are in contracts for water?
6. Is all water suitable for irrigation?
7. May water be supplied for irrigation by pumping?

**Methods of Applying Irrigation Water.** Choice of the various methods of applying irrigation water, of which there are about a dozen, is influenced by a consideration of (1) seasonal rainfall, (2) slope and general nature of the soil surface, (3) supply of water and how it is delivered, (4) crop rotation, and (5) permeability to water

of the soil and subsoil. A brief discussion of a few of the chief methods is given.

The *furrow method* frequently is used for the irrigation of fruit trees and row crops, such as potatoes, sugar beets, and corn or the grain sorghums. Furrows are made across the field, leading down the slope.



FIG. 92. Showing a method of distributing water from a flume into irrigation furrows. [From *California Agr. Ext. Service Circ. 16*, by M. R. Huberty and J. B. Brown.]

Water is let into the upper end of the furrow from a "head ditch" or pipe line running across the high end of the field.

The *corrugation system* is a modification of the furrow method and is used largely in the northwestern states for the irrigation of small-grain and hay crops because of the uneven topography, use of small streams of water, and prevailing methods of planting and harvesting. The irrigation furrows are placed at such distances apart and are of such length that will provide for the wetting of the soil between them by horizontal seepage. The permeability of the soil, slope of the land, and volume of water available are determining factors in spacing the furrows.

Irrigation by the *border method* is suitable on land having a slope in one direction of 1 inch to 2 feet per 100 feet, although the system has been used on land with much steeper slopes. The field is laid out in strips varying in width from 6 to 60 feet and in length according to the head of water available, the nature of the soil, and the degree of slope. If the slope of the field is too great, the strips may be laid out



FIG. 93. Irrigation by the corrugation method. Four corrugations are being supplied from one outlet. [From U.S.D.A. Farmers' Bull. 1348, by James C. Marr.]

diagonally across it; otherwise they run down the slope. The strips must be of such width and length that they may be entirely covered with water without letting the water in at a rate that would cause erosion of the soil or without permitting too much or too little water to soak into the soil at one end before it reaches the other end.

A simple formula has been proposed for figuring the proper area of the strips. It is based on two factors: (1) the head of water in cubic feet per second, and (2) the nature of the soil. For sandy soils the ratio of irrigating head to acres of land is 20 to 1, and for clay soils 2 to 1. Thus, with a flow of water of 10 cubic feet per second, the size of the strips would be 0.5 acre on sandy soil and 5.0 acres on clay soil.

The strips are set off by low, broad, flat-topped mounds of earth from 5 to 12 inches high, known as levees or borders. The borders are made in this shape so that farm implements may cross them readily and so that grazing stock will not destroy them when the field is in pasture. Needless to say, the soil surface in each strip must be leveled so that water will spread across it readily when let in through the two or more headgates (depending on the width of the strip) provided in the head ditch. It is not necessary that the strips in a field be at the same level.

The earliest and simplest method of irrigation is by *flooding*. This method is applicable principally to rather smooth land with a regular and moderate slope and when there is an abundant supply of water. When the fall is slight, the water is carried in shallow ditches from 50 to 100 feet apart in the direction of the slope. With a greater slope the ditches are made to wind across the field approximately on the contour. By making little earth dams in the ditches, the water is made to overflow and cover the whole field.

In fields with irregular surface or with a considerable fall, ridges called checks are thrown up on the contour, thus dividing the land into plots. Irrigation is generally started with the plots highest up the slope. When one plot has been irrigated, water is let into the next lower one from the ditch. It is not considered good practice to let the water from one plot into the next except in irrigating rice.

Another modification of the flooding system of irrigation is known as the *basin method*. The procedure consists in forming small basins by throwing up small earth dams or levees which retain the water while it soaks into the soil. A ditch is provided between each two rows of basins, and the water is run from the ditch into first one basin and then another as the irrigation proceeds. This system frequently is used in orchards and vineyards.

**Quantity of Water to Apply.** An ideal application of water would be a sufficient quantity to bring the soil, to the depth of the root zone of the crop, up to its field capacity. More water may result in the waterlogging of a portion of the subsoil or in the loss of water by drainage. If much water percolates below the root zone, it may accumulate under low areas, thus raising the water table unless suitable drainage facilities are provided. On the other hand, unless enough water is applied to result in appreciable drainage, it is difficult to remove excess salts from soils in which there is a tendency for alkali to accumulate.

The quantity of water available to plants which may be stored in soils ranges from  $\frac{1}{2}$  to 2 acre-inches per foot of depth. If the water-holding capacity of the soil per foot in depth is known, the approximate quan-

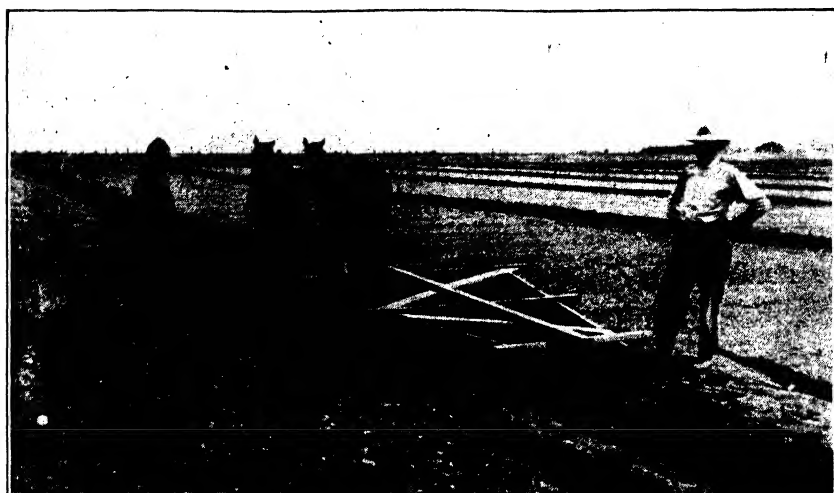


FIG. 94. A method of preparing land for the border method of irrigation.  
[From *U.S.D.A. Farmers' Bull.* 1243, by Samuel Fortier.]



FIG. 95. The irrigation of a peach orchard by the contour-check method.  
[From *California Agr. Ext. Service Circ.* 73, by J. B. Brown.]

tity of water to apply may be obtained by multiplying this figure by the depth in feet of root penetration of the crop. The readiness with which soils absorb water influences the quantity and rate of application. Most dry soils absorb water rapidly. Little change in the rate of water intake occurs in very permeable soils as the depth of wetting increases. Some clay soils, however, absorb water very slowly after the first foot or so has



FIG. 96. The basin method of irrigation. [From *U.S.D.A. Farmers' Bull.* 1518, by Samuel Fortier.]

become saturated. Because of this situation, it is advisable to hold the water for a long time on heavy soils and for only a short time on coarse-textured soils.

**Planning the Irrigation System.** Before deciding on the location, capacity, and design of irrigation and waste ditches and on the size of strips if this method of application is to be used, it is well to study the lay of the land carefully. This may be done by dividing the land into 100-foot squares and taking the elevation of each corner of the squares. Physical features, such as the location of streams, ditches, buildings, and roads, should also be shown on the map. By use of this map various arrangements of ditches and methods of applying water may be tried out on paper; thus is avoided the expense of staking them out to determine if they are feasible. When a layout is finally selected, it is well

to use temporary ditches and earth or canvas dams for the first year, to be sure the arrangement is satisfactory, before putting in permanent structures.

**Measurement of Irrigation Water.** The two kinds of units used in measuring irrigation water are (1) units of volume and (2) units of flow. Flow refers to the volume of water passing a given point in a unit of time. For example, the cross-sectional area in square feet of a stream or ditch, multiplied by the velocity of flow in feet per second, gives the cubic feet per second. The commonly used units of flow are cubic feet per second, gallons per minute, and miner's inch. The miner's

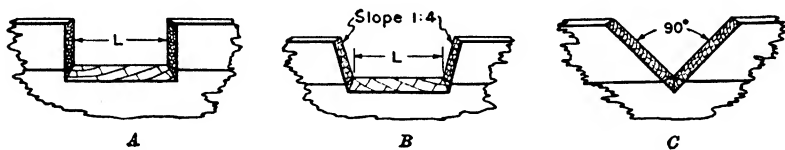


FIG. 97. Types of contracted weir notches: *A*, rectangular; *B*, trapezoidal; *C*, V-notch or 90-degree triangular. [From *California Agr. Exp. Sta. Bull.* 588, by J. E. Christiansen.]

inch is the flow through an orifice of 1 square inch under a specified "head," which usually varies from 4 to 6 inches. The head is the depth of water on the upstream side to the center of the orifice. It requires 38.5 to 50 miner's inches to equal a cubic foot per second, according to the regulations of different states which specify the height of the head of water.

Units of volume in common use are cubic foot, gallon, acre-foot, and acre-inch. The volume of water required to cover an acre to a depth of 1 inch is designated as an acre-inch.

For measuring the flow of small streams and ditches, various devices such as weirs, submerged orifices, Parshall measuring flumes,<sup>2</sup> and current meters are used. Many types of submerged orifices are in use, including the miner's-inch boxes. Two types of current meters are illustrated in Fig. 98. A wall or bulkhead across a ditch, with an opening cut in the top of it to permit water to flow through, is known as a weir. The opening may be rectangular, trapezoidal, or triangular and

<sup>2</sup> The Parshall measuring flume was developed through cooperative research by the U. S. Department of Agriculture and the Colorado Agricultural Experiment Station. It is named after R. L. Parshall, the engineer who conducted the experimental work. Water is measured by this flume much as it is by a weir. A full description of the device cannot be given here.

is called the weir notch. The height of the water above the bottom of the notch and the size and shape of the notch determine the quantity of water which will flow through it in a given time. The weir is one of the simplest and most commonly used means of measuring water.

**Contracts for Delivery of Water.** Agreements vary greatly concerning the quantity of water which is to be delivered to the land. A common method is to designate a stream of a given size, as 1 cubic foot per second for each 80 acres, with the provision that the purchaser will turn off the water when it is not needed. Sufficient water to cover the

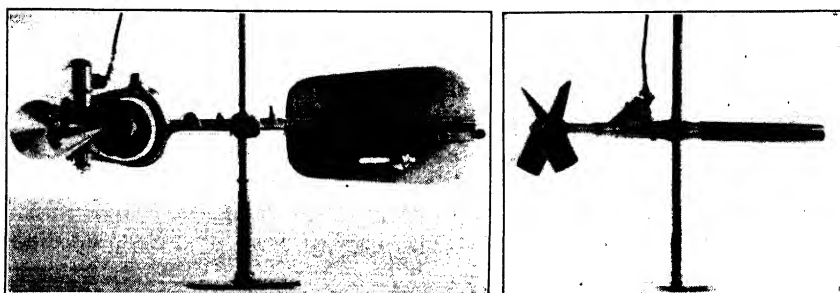


FIG. 98. Types of current meters with rod mountings. Left, Price type; right, Hoff type. Rod mountings are exchangeable for cable suspension. [From *California Agr. Exp. Sta. Bull. 588*, by J. E. Christiansen.]

land to a given depth, 2 feet during the season for example, or, in other words, 24 acre-inches, is frequently contracted for. Most contracts carry the provision that in case of water shortage the available supply will be prorated among the purchasers.

**Quality of Water.<sup>3</sup>** Careful attention should be given to the nature of the water to be used before an irrigation system is constructed. Sometimes the available water carries so high a concentration of soluble salts that its use for irrigation is not advisable, particularly on land which already contains a considerable concentration of alkali. Sodium salts in the water are much more objectionable than are salts of calcium and magnesium because of the tendency for sodium to cause deflocculation of the colloidal fraction of the soil and so develop an undesirable structure. Some waters also contain sufficient boron to prove toxic to plants with continued use. To make an estimate of the quality of irrigation water, the measurement of the following three characteristics is essential: (1) total concentration of salts, (2) the per cent of sodium,

<sup>3</sup> Much of the material dealing with quality of water was taken from *U.S.D.A. Technical Bull. 962*, by L. V. Wilcox.

and (3) the concentration of boron. The total concentration may be expressed as electrical conductivity, total equivalents per million of either anions or cations, or as dissolved solids.

In making an estimate of the quality of water the effect of the salts on both the soil and the plant must be taken into consideration. Various factors such as drainage, soil texture, and kind of clay minerals present

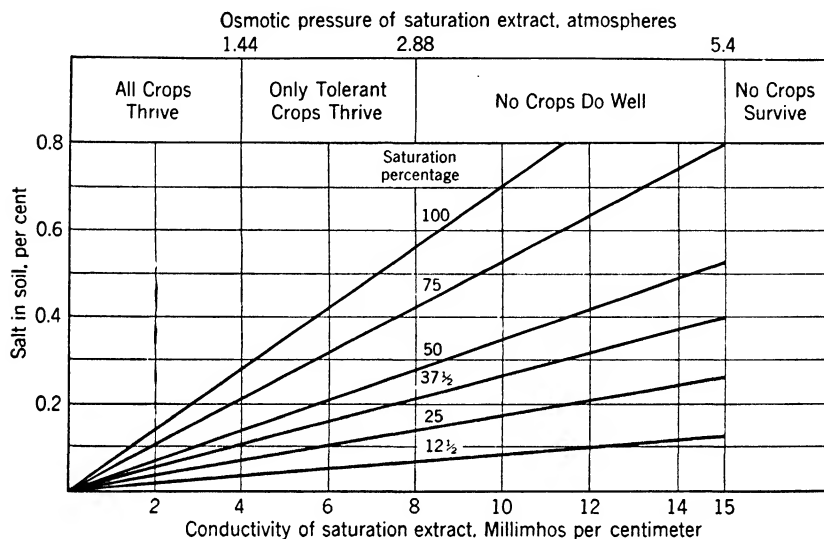


FIG. 99. A diagram for use in the interpretation of the analysis of an irrigation water. [From *Technical Bull. 962*, by L. V. Wilcox, Bureau of Plant Industry, Soils, and Agricultural Engineering of U.S.D.A.]

influence the effects on the soil. The ultimate effect on the plant is the result of these two and other factors operating simultaneously. Consequently, there can be no method of interpretation which is absolutely accurate under all conditions. The schemes for interpretation are accordingly based chiefly on experience. The classification of waters set forth in Table 48 and suggested by Scofield has been found reasonably satisfactory. Scofield has also proposed the limits of boron content for different groups of crops which are shown in Table 49. Some workers think that a somewhat better classification of waters, especially in regard to sodium content, is that presented in Fig. 100. By using total concentration in equivalents per million and per cent of sodium, a point may be located on the chart which falls in one of the areas of designated water quality.

## Irrigation

TABLE 48

PERMISSIBLE LIMITS FOR ELECTRICAL CONDUCTIVITY AND PER CENT OF SODIUM OF SEVERAL CLASSES OF IRRIGATION WATER

| Classes of Water |             | Electrical Conductivity             | Sodium, per cent |
|------------------|-------------|-------------------------------------|------------------|
| Rating           | Grade       |                                     |                  |
| 1                | Excellent   | E.C. $\times 10^6$ at 25°C.<br><250 | <20              |
| 2                | Good        | 250–750                             | 20–40            |
| 3                | Permissible | 750–2,000                           | 40–60            |
| 4                | Doubtful    | 2,000–3,000                         | 60–80            |
| 5                | Unsuitable  | >3,000                              | >80              |

TABLE 49

PERMISSIBLE LIMITS FOR BORON OF SEVERAL CLASSES OF IRRIGATION WATER

| Classes of Water |             | Sensitive Crops, p.p.m. | Semi-Tolerant Crops, p.p.m. | Tolerant Crops, p.p.m. |
|------------------|-------------|-------------------------|-----------------------------|------------------------|
| Rating           | Grade       |                         |                             |                        |
| 1                | Excellent   | <0.33                   | <0.67                       | <1.00                  |
| 2                | Good        | 0.33–0.67               | 0.67–1.33                   | 1.00–2.00              |
| 3                | Permissible | 0.67–1.00               | 1.33–2.00                   | 2.00–3.00              |
| 4                | Doubtful    | 1.00–1.25               | 2.00–2.50                   | 3.00–3.75              |
| 5                | Unsuitable  | >1.25                   | >2.50                       | >3.75                  |

**Irrigation by Pumping.** Irrigation by pumping has been increasing in popularity for many years. The system is used in arid regions in conjunction with irrigation from streams and reservoirs, in semi-arid areas to supplement the limited rainfall, and in humid climates for the production of crops of high value per acre, especially fruits, vegetables, and nursery stock.

After years of irrigation, the water table under some of the western reclamation projects became so high that it was necessary to install pumps at frequent intervals to lower it. The water so removed from the soil was used again for irrigation, thus supplementing the original

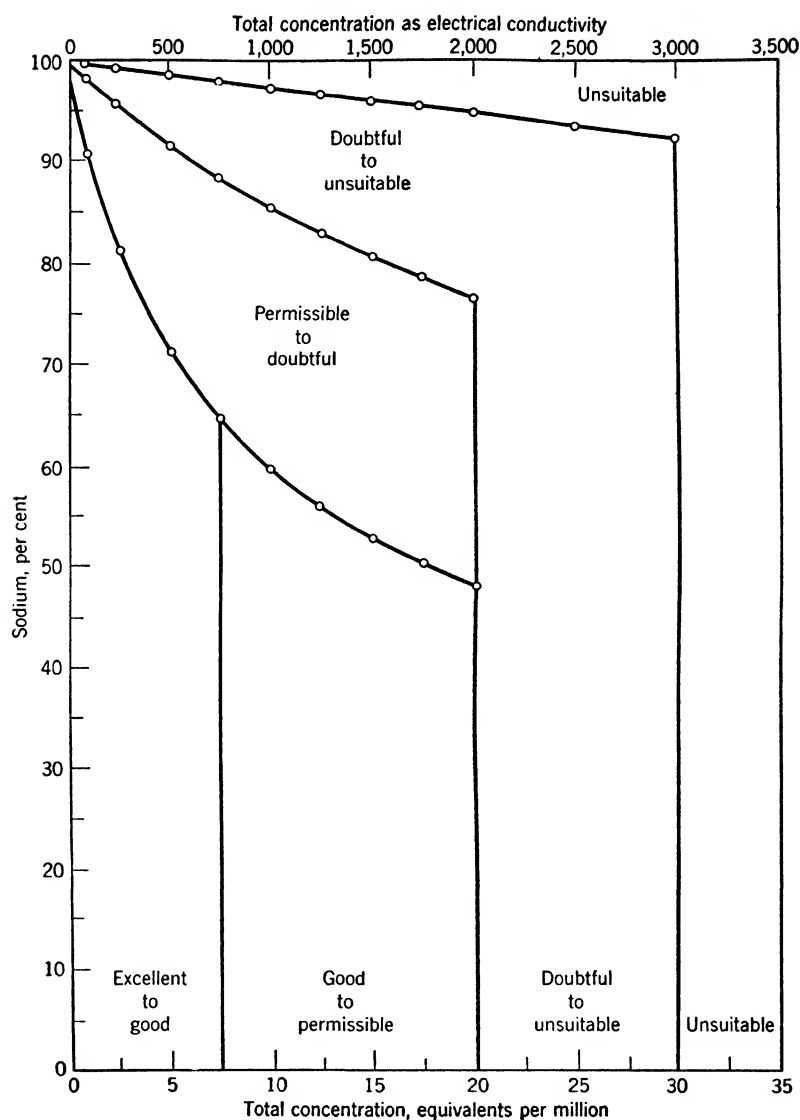


FIG. 100. A graphical representation of the relation between crop growth, conductivity, and osmotic pressure of the saturated extract and the per cent of sodium in saline soils. The statements in the enclosed areas of the diagram such as "permissible to doubtful," "doubtful to unsuitable," etc., refer to suitability for crop growth. [From "Diagnosis and Improvement of Saline and Alkali Soils," U. S. Regional Salinity Laboratory, Riverside, California.]

supply and providing for the watering of more land. Wells of great depth and about 24 inches in diameter have also been bored at the borders of irrigated valleys where the elevation is too high to permit obtaining water from canals.

In the Great Plains states large acreages are irrigated by pumping water from the underground flow of streams which usually sink below the surface of the previous soil except during flood periods.



FIG. 101. Irrigating sugar beets in Ohio with well water. The whole system with the exception of the well is portable. [From *U.S.D.A. Farmers' Bull.* 1846, by F. E. Staebner.]

The depth from which water can be lifted economically for irrigation depends largely on the quantity of water which will be needed and on the value of the crop to be grown as well as on the cost of the fuel consumed in operating the pumps.

Before a pumping system is installed, careful investigation and tests should be made to determine the constancy of the water supply, particularly in dry seasons, the quantity of water the well will deliver under constant pumping at the required rate to irrigate the desired acreage, and the depth to which the water is lowered in the well by pumping.

The water pumped from streams, ponds, and wells may be applied to the soil by any of the methods previously described. In addition,

overhead sprinkling systems are frequently used on small acreages. Lines of pipe carrying small nozzles every few feet are mounted on posts at a conventional height and are so arranged that the pipes rotate through an angle of somewhat less than 180°. A system which has been increasing in favor rapidly makes use of rotating sprinklers similar to those used on lawns and golf courses but larger. The development of light-weight pipe with self-locking joints has given great impetus to supplemental irrigation in the humid areas. These pipes may be moved readily from one part of the field to another; thus they permit the irrigation of a considerable area with a minimum of equipment.

Improvement in equipment and comparatively high prices for vegetables and small fruits has resulted in a rapid increase in acreage of land irrigated in the eastern states. For example, in 1944 Massachusetts had 11,355, New Jersey 11,712, and Florida 221,917 acres under irrigation. The acreage of land under irrigation in the different states is shown in Table 50.

TABLE 50

## IRRIGATED LAND IN FARMS IN THE VARIOUS STATES, 1944

| State         | Acres<br>Irrigated | State          | Acres<br>Irrigated |
|---------------|--------------------|----------------|--------------------|
| Alabama       | 487                | Nebraska       | 631,762            |
| Arizona       | 736,027            | Nevada         | 674,204            |
| Arkansas      | 288,665            | New Hampshire  | 65                 |
| California    | 4,952,819          | New Jersey     | 11,712             |
| Colorado      | 2,698,579          | New Mexico     | 534,640            |
| Connecticut   | 496                | New York       | 10,316             |
| Delaware      | 22                 | North Carolina | 229                |
| Florida       | 221,917            | North Dakota   | 22,814             |
| Georgia       | 423                | Ohio           | 4,178              |
| Idaho         | 2,026,280          | Oklahoma       | 2,237              |
| Illinois      | 368                | Oregon         | 1,129,059          |
| Indiana       | 830                | Pennsylvania   | 8,764              |
| Iowa          | 1,197              | Rhode Island   | 133                |
| Kansas        | 96,248             | South Carolina | 62                 |
| Kentucky      | 250                | South Dakota   | 52,895             |
| Louisiana     | 535,619            | Tennessee      | 393                |
| Maine         | 100                | Texas          | 1,320,216          |
| Maryland      | 287                | Utah           | 1,124,081          |
| Massachusetts | 11,355             | Vermont        | 6                  |
| Michigan      | 2,850              | Virginia       | 1,419              |
| Minnesota     | 210                | Washington     | 520,153            |
| Mississippi   | 3                  | West Virginia  | 42                 |
| Missouri      | 1,113              | Wisconsin      | 4,569              |
| Montana       | 1,555,480          | Wyoming        | 1,353,875          |

# 20

## Soil Resources

As food, textiles, and housing materials are largely products of the soil, either directly or indirectly, an ample acreage of productive land is one of the most essential and stabilizing resources a people can have. A nation with soil resources so limited that it must depend mainly on imported food and fiber is always in a precarious economic position because not only war but also changes in trade conditions may curtail the supply of imports. Furthermore, manufacturers of many commodities must depend on agricultural workers to purchase a considerable portion of the manufactured products, even though the workers live in different countries. Unless the people who live on the land have reasonable incomes, their purchasing power is limited. It is highly important, therefore, to the city producer that there be ample good land for farming purposes and that the land be maintained in a sufficiently productive state to afford a reasonable income to farmers and to provide an adequate food supply.

In considering the soil resources of the United States and other countries, and the adequacy of these resources to supply the needs of the peoples concerned, the following major topics should be investigated.

### Objectives

- A. Acreage of producing and potential farm land in the United States.
- B. Acreages of arable land in different countries, and land requirement per capita to supply food.
- C. Land policies of the United States.

### ACREAGE OF FARM LAND IN THE UNITED STATES

An inventory of land resources in continental United States was made by federal and state authorities. Liberal use was made of data contained in the 1935 Census of Agriculture. Objectives of the survey were, first, to determine the acreage of land suitable for cultivation without erosion

damage under prevalent farming methods and with price levels equal to those prevailing during the period 1921 to 1936; secondly, to know the additional acreage that could be profitably farmed under the same price levels if the best farming methods and erosion control practices were employed. Other studies have been made to determine the acreage of first-, second-, and third-class land in our country and to indicate the proportion of the various classes of land in the different states. Answers to the following questions may well be sought in considering this topic.

### Questions

1. How much land is suitable for cultivation without special provisions to control erosion?
2. To what uses is this land being put at present?
3. Where is this good land located?
4. How much land can be farmed if soil-conserving methods are used?
5. What is the total acreage of present and potential crop land?
6. What is the distribution of land of different quality among the states?

**Land Suitable for Cultivation.** The 1935 Census of Agriculture listed 415,334,931 acres as the crop-land area of continental United States. This figure includes the acreage of (1) crops harvested, (2) land on which crops failed, and (3) land suitable for cropping but temporarily idle or fallow. The best estimates indicate that only 39.1 per cent of this crop land, amounting to 160,948,703 acres, is of a quality that can be cultivated by prevailing methods, without serious erosion, and yield well enough to give profitable returns if prices are as they were between 1921 and 1936. This means either that 60.9 per cent of the present crop land is of such poor quality that it cannot be profitably farmed or that it is subject to continued erosion under the methods by which it is being cultivated. Under price levels which have prevailed during and after World War II a larger percentage of this land doubtless could be farmed profitably.

**Present Use of Good Land.** All the land which may be profitably farmed without serious erosion is not now listed in the United States census as crop land. There are some 21,611,680 acres, designated as plowable land, which are now pasture. Brush or timber occupies 19,856,239 acres. Also, 6,429,565 acres need drainage, and 2,919,500 acres could be farmed if irrigated. Summing up these figures, we find that 50,816,984 acres can be added to the present area of crop land suitable for cultivation under present practices and at the price levels mentioned, provided that the needed improvements of clearing, drain-

age, irrigation, and the like, are made. This gives a total of 211,765,-687 acres of what may be called good land.

**Location of Potential Crop Land.** It is of interest to note where this good land, which may be brought into production if needed, is located. The location and the type of improvement needed are shown by the data in Table 51.

TABLE 51

POTENTIAL CROP LAND SUITABLE FOR CULTIVATION UNDER PREVAILING FARMING METHODS AND AT PRICE LEVELS SIMILAR TO THOSE OF 1921 TO 1936 \*

| Group of States    | Present Use or Improvement Needed |                          |                           |                             | Total      |
|--------------------|-----------------------------------|--------------------------|---------------------------|-----------------------------|------------|
|                    | Acres in plowable pasture         | Acres in brush or timber | Acres in need of drainage | Acres in need of irrigation |            |
| New England        | 103,143                           | .....                    | .....                     | .....                       | 103,143    |
| Middle Atlantic    | 369,129                           | 22,843                   | 1,984                     | .....                       | 393,956    |
| South Atlantic     | 1,015,910                         | 4,003,802                | 1,873,609                 | 600                         | 6,893,921  |
| East North Central | 6,568,996                         | 3,127,172                | 678,496                   | .....                       | 10,374,664 |
| West North Central | 7,094,306                         | 285,923                  | 1,191,600                 | 138,100                     | 8,709,929  |
| East South Central | 1,502,272                         | 3,790,762                | 222,021                   | .....                       | 5,515,055  |
| West South Central | 3,347,079                         | 8,238,937                | 1,994,323                 | 790,706                     | 14,371,045 |
| Pacific            | 417,900                           | 274,600                  | 205,000                   | 236,000                     | 1,133,500  |
| Mountain           | 1,192,945                         | 112,200                  | 262,532                   | 1,754,094                   | 3,321,771  |
| Total              | 21,611,680                        | 19,856,239               | 6,429,565                 | 2,919,500                   | 50,816,984 |

\* Compiled from data in "The Problem: The Nation As a Whole," E. J. Utz, C. E. Kellogg, E. H. Reed, J. H. Stallings, and E. N. Munns, *U.S.D.A. Yearbook of Agriculture* 1938, p. 95.

Plowable pasture offers the greatest opportunity for expanding our present acreage of good land for producing crops; clearing off brush and timber, drainage, and irrigation, in the order named, are the other possibilities. The west South Central States have the greatest acreage which may be brought into production. More than half of this potential crop land is in timber or brush. The east North Central States have the second largest acreage of good land which is not under cultivation. Considerably more than half of this land is in pasture. The west North Central and South Atlantic states follow in order in area of land suitable for crop production if needed improvements are made.

**Land Suitable for Cultivation by Use of Soil-Conserving Methods.** To the 39.1 per cent of the "crop land" which is considered suitable for cultivation under present methods may be added 42.9 per cent, or 178,178,685 acres, which may be farmed profitably if soil-conserving methods are used. Soil-conserving methods of farming are discussed in Chapter 16. This leaves 18 per cent of the present land listed in the census as crop land which is considered either too poor to yield profitable returns at prices which prevailed between 1921 and 1936 or else of such rough topography that its fertility cannot be maintained even by the use of good soil-conserving practices.

In addition to the 82 per cent (39.1 per cent + 42.9 per cent), 339,079,482 acres, of the present crop land which may be farmed by present or improved methods, there can be added a considerable area which needs clearing, drainage, irrigation, or plowing before it can be farmed. Furthermore, liberal applications of lime and fertilizer would be required on a large area of the land. Soil-conserving methods of farming would then be needed if the land were to be cropped profitably for any length of time. The location of these lands and the improvement needed are shown in Table 52. The table also gives the acres of land which are now in cultivation and on which soil-conserving methods should be used.

The west North Central States have by far the largest area of land which may be cropped profitably by use of soil-conserving methods, and incidentally about three-fourths of this land is now being cultivated regardless of the extent to which soil-conserving methods are being used. The west South Central States have the next largest acreage of land that falls into the class under discussion, with slightly more than one-half of the land being cultivated. The east North Central States and the South Atlantic States offer about equal opportunities for enlarging the acreage of crop land by use of soil-conserving farming methods. Approximately 62 per cent of the land which should be cultivated only by special methods to maintain its productivity is already under cultivation. This situation is deserving of careful consideration, for the question immediately arises, how long will these soils remain in the productive class under present methods of management?

Plowable pasture again occupies the greatest acreage which may be brought into cultivation, if need be, and again the clearing of brush and timber offers the second largest opportunity to increase crop acreage.

**Total Acreage of Present and Potential Crop Land.** Summing up the data presented in the preceding paragraphs shows that there are some 160,948,703 acres of land in cultivation in the United States which

TABLE 52

LAND SUITABLE FOR CULTIVATION IF SOIL-CONSERVING METHODS ARE USED AND IF THE INDICATED IMPROVEMENTS ARE MADE \*

| Group of States    | Present Use or Improvement Needed |                          |                           |                             |             |                      |             |
|--------------------|-----------------------------------|--------------------------|---------------------------|-----------------------------|-------------|----------------------|-------------|
|                    | Acres in plowable pasture         | Acres in brush or timber | Acres in need of drainage | Acres in need of irrigation | Total       | Acres in cultivation | Total       |
| New England        | 336,127                           | .....                    | .....                     | .....                       | 336,127     | 1,842,496            | 2,178,623   |
| Middle Atlantic    | 1,620,920                         | 73,276                   | 1,984                     | .....                       | 1,696,180   | 10,016,169           | 11,712,349  |
| South Atlantic     | 3,419,817                         | 11,552,831               | 2,977,376                 | 600                         | 17,950,624  | 18,265,787           | 36,216,411  |
| East North Central | 12,313,011                        | 5,878,757                | 690,611                   | .....                       | 18,882,379  | 19,337,066           | 38,219,445  |
| West North Central | 20,248,118                        | 599,393                  | 1,213,900                 | 349,800                     | 22,411,211  | 68,475,596           | 90,886,807  |
| East South Central | 5,138,369                         | 7,203,114                | 222,021                   | .....                       | 12,563,504  | 9,923,926            | 22,487,430  |
| West South Central | 5,986,678                         | 15,704,103               | 2,025,803                 | 872,671                     | 24,589,255  | 30,058,149           | 54,647,404  |
| Pacific Mountain   | 1,162,500                         | 958,400                  | 239,700                   | 2,087,000                   | 4,447,600   | 10,091,122           | 14,538,722  |
| Total              | 52,722,676                        | 42,151,544               | 7,663,821                 | 5,848,729                   | 108,386,770 | 178,130,779          | 286,517,549 |

\* Compiled from data in "The Problem: The Nation As a Whole," E. J. Utz, C. E. Kellogg, E. H. Reed, J. H. Stallings, and E. N. Munns, *U.S.D.A. Yearbook of Agriculture*, 1938, p. 95.

may be profitably farmed with present methods without serious deterioration through erosion. To the above, 50,816,984 acres may be added by plowing up pasture, clearing, draining, and irrigating. There is a total, then, of 211,765,687 acres suitable for profitable farming with present methods. If soil-conserving methods were universally adopted and the needed improvements made, this area could be enlarged by 286,517,549 acres. A considerable area of this land is now being cultivated regardless of the damage to the soil. This gives a grand total of 498,283,236 acres which may be used with reasonable profit to the farmer for the production of food, hay, and fiber crops if conditions demand and if soil-conserving methods are used.

**Distribution of Land of Different Quality among the States.** A study has been made, with the purpose of dividing the land into five groups, on the basis of the land's capacity to produce grains, grasses, and legumes without irrigation, drainage, or fertilization. The group names used are excellent, good, fair, poor, and unfit for crop production. In Table 53 is presented the total percentage of land in the different groups and the percentage of land in each class found in each of

# Distribution of Land of Different Quality

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TABLE 53

**APPROXIMATE ACREAGE OF LAND; EXCELLENT, GOOD, FAIR, POOR, OR UNFIT FOR GENERAL CROPS WITHOUT IRRIGATION, DRAINAGE, OR FERTILIZATION IN THE SEVERAL STATES AND IN CONTINENTAL UNITED STATES \***

| Quality Rating  | Excellent           | Good                             | Fair  | Poor                   | Unfit   |
|-----------------|---------------------|----------------------------------|---|------------------------|---|
| % of total land | 5.3%                | 11.1%                            | 18.1%   | 19.1%                  | 46.4%   |
| United States   | 101,038,000         | 210,935,000                      | 345,872,000   | 362,559,000            | 881,735,000   |
| Iowa            | 25,983,000          | 6,906,000                        | 1,393,000   | 1,007,000              | 345,000   |
| Illinois        | 14,777,000          | 6,847,000                        | 6,223,000   | 6,622,000              | 961,000   |
| Minnesota       | 12,022,000          | 12,139,000                       | 7,511,000   | 6,899,000              | 13,145,000  |
| Missouri        | 8,075,000           | 13,833,000                       | 12,304,000  | 4,259,000              | 4,915,000   |
| Nebraska        | 8,121,000           | 9,690,000                        | 9,640,000   | 10,908,000             | 10,758,000  |
| Indiana         | 5,262,000           | 6,438,000                        | 7,744,000   | 2,437,000              | 1,153,000   |
| Ohio            | 4,214,000           | 6,234,000                        | 10,440,000  | 3,810,000              | 1,375,000   |
| Kansas          | 3,765,000           | 15,172,000                       | 15,965,000  | 11,207,000             | 6,095,000   |
| South Dakota    | 3,052,000           | 8,775,000                        | 8,960,000   | 15,552,000             | 12,821,000  |
| Wisconsin       | 2,820,000           | 14,530,000                       | 8,397,000   | 4,564,000              | 5,271,000   |
| Michigan        | 2,251,000           | 8,961,000                        | 5,387,000   | 7,229,000              | 13,223,000  |
| Oklahoma        | 1,701,000           | 12,795,000                       | 15,269,000  | 7,826,000              | 6,856,000   |
| Texas           | 1,591,000           | 19,461,000                       | 54,974,000  | 35,950,000             | 56,098,000  |
| Arkansas        | 1,453,000           | 7,952,000                        | 9,231,000   | 8,300,000              | 6,707,000   |
| Louisiana       | 1,289,000           | 2,973,000                        | 7,347,000   | 12,443,000             | 5,024,000   |
| Tennessee       | 902,000             | 4,770,000                        | 9,305,000   | 8,240,000              | 3,412,000   |
| Kentucky        | 864,000             | 5,020,000                        | 8,956,000   | 7,906,000              | 2,877,000   |
| Mississippi     | 844,000             | 5,336,000                        | 8,615,000   | 10,929,000             | 4,091,000   |
| West Virginia   | 389,000             | 1,350,000                        | 2,987,000   | 7,647,000              | 2,828,000   |
| Oregon          | 343,000             | 2,059,000                        | 4,694,000   | 5,465,000              | 48,568,000  |
| Pennsylvania    | 177,000             | 6,268,000                        | 9,497,000   | 5,909,000              | 6,909,000   |
| Maryland        | 144,000             | 1,875,000                        | 1,446,000   | 1,839,000              | 973,000   |
| New York        | 94,000              | 6,864,000                        | 9,249,000   | 9,172,000              | 4,926,000   |
| Washington      | 84,000              | 542,000                          | 6,537,000   | 6,760,000              | 28,836,000  |
| Florida         | 71,000              | 933,000                          | 3,928,000   | 12,893,000             | 17,078,000  |
| Connecticut     | 48,000              | 339,000                          | 902,000   | 962,000                | 851,000   |
| California      | 47,000              | 478,000                          | 3,365,000   | 13,380,000             | 82,347,000  |
| Vermont         | 33,000              | 901,000                          | 1,268,000   | 2,185,000              | 1,452,000   |
| Massachusetts   | 12,000              | 760,000                          | 1,275,000   | 1,644,000              | 1,367,000   |
| New Jersey      | 6,000               | 943,000                          | 964,000   | 1,379,000              | 1,471,000   |
| New Hampshire   | 2,000               | 237,000                          | 370,000   | 1,672,000              | 3,478,000   |
| North Dakota    | .....               | 7,366,000                        | 17,716,000  | 14,152,000             | 5,680,000   |
| Virginia        | .....               | 3,653,000                        | 9,280,000   | 7,312,000              | 5,400,000   |
| Alabama         | .....               | 3,002,000                        | 9,528,000   | 10,730,000             | 9,491,000   |
| Georgia         | .....               | 1,852,000                        | 15,614,000  | 9,750,000              | 10,369,000  |
| Maine           | .....               | 1,615,000                        | 3,789,000   | 5,791,000              | 7,978,000   |
| North Carolina  | .....               | 1,167,000                        | 11,360,000  | 9,382,000              | 9,285,000   |
| South Carolina  | .....               | 296,000                          | 7,070,000   | 7,128,000              | 5,023,000   |
| Colorado        | .....               | 286,000                          | 7,041,000   | 13,744,000             | 45,270,000  |
| Delaware        | .....               | 222,000                          | 351,000   | 417,000                | 273,000   |
| Idaho           | .....               | 87,000                           | 949,000   | 3,729,000              | 48,581,000  |
| Rhode Island    | .....               | 5,000                            | 228,000   | 71,000                 | 378,000   |
| Distribution    | ¾ in corn belt area | ½ between Texas and North Dakota | ½ south of Potomac and Ohio Rivers, including Arkansas, Louisiana, Oklahoma and Texas | 43% in southern states | 75% of all U.S. Non-tillable land is in the 11 far western states owing to lack of rain |

\* "A Graphic Summary of Physical Features and Land Utilization in the United States," O. E. Baker, *U.S.D.A. Misc. Pub.* 260, 1937, pp. 12-14.

the forty-two states having excellent or good land. The distribution of the first three classes of land is also shown graphically in Fig. 102.

The high percentage of excellent and good land in the Corn Belt states and the states immediately adjoining them is worthy of consideration. The question may well be asked, what conditions gave rise to the development of so much land of superior quality in that location? It should be remembered, furthermore, that some of the land designated "fair" in this classification produces superior yields when fertilized. In fact, some of the states which have very little if any "excellent" land have higher average acre yields of several of the grain crops than do the Corn Belt states. Likewise, the production on large acreages of irrigated land is much greater than on high-quality land without irrigation.

A more recent study has been made by the Soil Conservation Service of the suitability of the land now in farms for use under different methods of management. The results are presented in Table 54. These figures show that only 460,000,000 acres of the 1,142,000,000 acres in farms, or about 40 per cent, are suitable for regular cultivation. Furthermore, about 48 per cent of this land requires intensive treatment before it can be safely farmed. There are 554,000,000 acres, or approximately 48 per cent of the total land in farms, which is not suited for cultivation but should be used for grazing or forestry.

#### **ACREAGES OF ARABLE LAND AND LAND REQUIREMENTS**

A knowledge of the acreage of farm land in different countries in relation to population leads to a better understanding of some of the problems confronting the great nations. The interdependence of nations is brought out also by such a study. The more one learns about good farm land and the part it plays in the welfare of people, the more respect one has for the soil. Answers to the following questions will assist in obtaining an understanding of the land problem in different countries.

#### **Questions**

1. What is meant by arable land?
2. What fraction of the earth's surface is arable?
3. How much arable land per capita is there in different countries?
4. Are American diets well balanced?
5. What is the land requirement per capita to supply food?
6. What increase in food production will be needed by 1960?

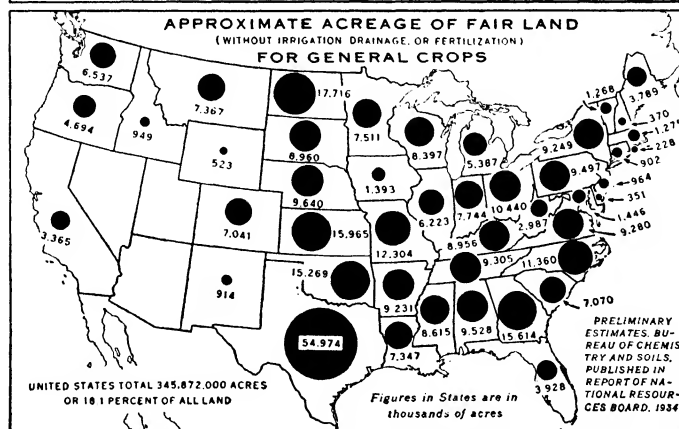
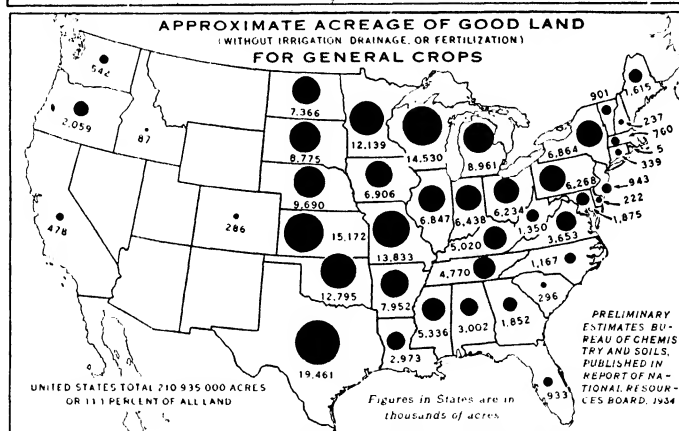
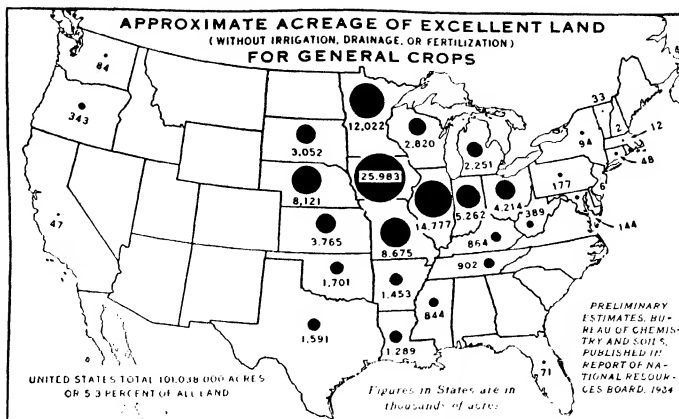


FIG. 102. The distribution of excellent, good, and fair land in the various states. [From U.S.D.A. *Misc. Pub.* 260.]

TABLE 54

DISTRIBUTION OF LAND IN FARMS IN THE UNITED STATES ACCORDING  
TO CAPABILITY \*

|  | Millions<br>of Acres<br>(approx.) † | Millions<br>of Acres<br>(approx.) |
|--|-------------------------------------|-----------------------------------|
| Suited for regular cultivation   |                                     | 460                               |
| Class I: Very good land; few or no limitations; can be cultivated safely with ordinary good farming methods  | 70                                  |                                   |
| Class II: Good land; moderate limitations or hazards; can be cultivated safely with moderately intensive treatments  | 170                                 |                                   |
| Class III: Moderately good land; severe limitations or hazards; can be cultivated safely with intensive treatments   | 220                                 |                                   |
| Suited for limited cultivation   |                                     | 95                                |
| Class IV: Fairly good land; very severe limitations or hazards; suited for some forms of limited cultivation or for limited choice of crops but usually best suited for pasture or hay | 95                                  |                                   |
| Not suited for cultivation but suited for grazing or forestry  |                                     | 554                               |
| Class V: Not more than slight limitations  | 40                                  |                                   |
| Class VI: Moderate limitations   | 250                                 |                                   |
| Class VII: Severe limitations  | 264                                 |                                   |
| Not suited for cultivation, grazing, or forestry   |                                     | 33                                |
| Class VIII: Suited for wildlife or recreational use, or for watershed protection purposes  | 33                                  |                                   |
| Total land in farms reported by the 1945 Census of Agriculture   |                                     | 1,142                             |

\* From Proceedings of the Inter-American Conference on Conservation of Renewable Natural Resources, Denver, Colorado, Sept. 7-20, 1948.

† Inasmuch as these estimates are based on surveys of only limited sample areas, they are subject to revision as more land is surveyed.

**Meaning of Arable.** A broad and variable interpretation of the term arable is possible on the basis of experience and viewpoint. Strictly speaking, arable land is land which is suitable for plowing or cultivation. No mention is made, however, of whether such land may be cultivated profitably at any set price level for agricultural products or whether the land will deteriorate rapidly under cultivation unless special tillage practices and cropping systems are used. Probably there is a tendency to apply the term arable from a purely physical viewpoint and to consider any cleared land arable which is not too stony, too rough topograph-

ically, too badly in need of drainage to grow crops, or which is in an area of very limited rainfall during the growing season. It is evident that land which one person or group of persons would consider arable might not be so classed by another group of persons. For this reason it is difficult to compare accurately the acreages of good land in different countries. In the following discussion the comparisons are based on the best data available.

**About One-Tenth of the World's Land in Cultivation.** In general, about one-half of the land in the world is considered entirely unsuitable for cultivation. This portion is made up of areas permanently covered with ice and snow (11 per cent), the tundra (4 per cent), high mountains (16 per cent), and deserts and semi-deserts (17 per cent). Although appreciable areas of desert may be irrigated, they are not large enough to change the general picture. By no means all of the other one-half of the world's land can be easily placed under cultivation. Much of it is so rocky, so sandy, so hilly, so full of soluble salts, or so badly in need of drainage that it may never be cultivated. Estimates of the percentage of the world's land now in cultivation vary widely, some being greater and some less than 10 per cent. Certainly not a large part of the land surface is being cultivated. How much this area can be increased practically depends on economic conditions and scientific developments, but any large increase in the immediate future appears doubtful.

**Acreage of Arable Land per Capita.** If the land designated in the census as crop land be considered arable land, continental United States has 415,334,931 acres of arable land. Assuming a population of 150,000,000 persons, this allows approximately 2.77 acres per capita. The preceding discussion, however, shows that much of the crop land is being farmed without profit or at the expense of serious deterioration through erosion. It has been brought out that, by improving the good land not now in crops, a total of 211,765,687 acres could be made available which may be profitably farmed by present methods without serious damage by erosion. There are about 1.41 acres of such land per person. By bringing into cultivation all land which may be profitably farmed, without severe erosion, by use of soil-conserving methods there would be 498,283,236 acres, or 3.32 acres per capita.

By making use of available data, Table 55 has been prepared to give a picture of the land per capita available for crop production in several countries. These data show in how favorable a position the United States finds itself with regard to land resources. The precarious position of several countries with respect to soils on which to produce food

TABLE 55

POPULATION, ARABLE LAND, AND ARABLE LAND PER CAPITA IN  
SEVERAL COUNTRIES \*

| Country                            | Population  | Arable Land   | Acres of Arable Land per Capita |
|------------------------------------|-------------|---------------|---------------------------------|
| Belgium                            | 8,257,400   | 2,527,833     | 0.31                            |
| Canada                             | 11,385,000  | 59,517,120    | 5.23                            |
| China                              | 450,000,000 | 189,270,800 † | 0.40                            |
| Denmark                            | 3,844,300   | 6,609,720     | 1.72                            |
| England }<br>Wales }<br>Scotland } | 46,024,000  | 11,505,260    | 0.25                            |
| Finland                            | 3,887,200   | 6,439,290     | 1.66                            |
| France                             | 41,400,000  | 51,205,570    | 1.24                            |
| Germany }<br>Austria }             | 79,375,000  | 54,910,570    | 0.69                            |
| Netherlands                        | 9,007,700   | 2,556,450     | 0.28                            |
| Italy                              | 42,918,000  | 32,117,410    | 0.75                            |
| Japan                              | 69,254,000  | 14,889,160    | 0.22                            |
| U.S.S.R.                           | 169,519,000 | 553,072,520   | 3.26                            |
| India                              | 275,541,000 | 374,860,584   | 1.32                            |
| United States ‡                    | 150,000,000 | 415,334,931   | 2.77                            |

\* Sources of data: *The China Yearbook*, 1935-1936, pp. 123 and 839; *International Yearbook of Agricultural Statistics*, 1939-1940, Introduction, p. x; *Statistical Yearbook of League of Nations*, 1942-1943.

† Cultivated land. Total acreage is somewhat in excess of this figure because surveys are not complete.

‡ Estimate for 1950.

and clothing for their people is very evident. Perhaps this scarcity of soil resources is a contributing factor to the social unrest in certain countries and to activities which continually disturb world peace.

**American Diets in Need of Balancing.** Studies have been made of the diets of Americans living in towns and cities, on farms, and in villages and near cities. The results collected in 1936 showed that less than one-fifth of all families had diets that met the recommendations of the National Research Council for all the seven nutrients considered. This finding, of course, does not mean that all these people were hungry, although some did not have enough food. It does mean that the diets of many families were not properly balanced. Surveys in 1942 showed that consumption of food energy averaged higher in all income groups than that contained in planned, adequate diets. There were, however, appreciable shortages in the consumption of calcium, iron, vitamin A, ascorbic acid, niacin, thiamin, and riboflavin. It would appear, then,

that as a whole Americans need to exercise more care in the selection of their foods rather than to consume more food. In Table 56 are given two planned, adequate diets and also the allowance of various nutrients recommended by the National Research Council.

TABLE 56

ADEQUATE DIET PLANS AND NUTRIENT ALLOWANCES RECOMMENDED BY THE  
NATIONAL RESEARCH COUNCIL \*

| Food Nutrient | Unit               | Adequate Diet Plans |               | Recommended Allowance by NRC |
|---------------|--------------------|---------------------|---------------|------------------------------|
|               |                    | Low-cost            | Moderate-cost |                              |
| Food energy   | Calorie            | 2,720               | 2,810         | 2,640                        |
| Protein       | Gram               | 85                  | 92            | 65                           |
| Vitamin A     | International unit | 8,530               | 10,330        | 4,580                        |
| Calcium       | Milligram          | 1,080               | 1,250         | 940                          |
| Iron          | "                  | 15.0                | 15.9          | 11.7                         |
| Thiamin       | "                  | 1.8                 | 1.9           | 1.30                         |
| Riboflavin    | "                  | 2.2                 | 2.5           | 1.78                         |
| Niacin        | "                  | 17.0                | 17.8          | 13.0                         |
| Ascorbic acid | "                  | 128.0               | 159.0         | 71.0                         |

\* From Bureau of Human Nutrition and Home Economics. Nutrients are those contained in food as brought into the kitchen; no allowance is made for waste or losses in cooking.

**Land Required to Produce Food.** There arises the question whether there is enough good land in continental United States to produce the foods needed to supply adequate diets for all our people. To obtain an answer to this question we must know how many acres are required to produce the foods needed by one person. This point has been given careful consideration. Table 57 is quoted from "Efficient Use of Food Resources in the United States," by R. P. Christensen, *U. S. Department of Agriculture Technical Bulletin 963*.

In calculating the acres of crop land required per person, the acreage of pasture was converted into crop land equivalent. In the 2.67 acres required for the 1943-45 civilian diet, pasture accounts for about 0.9 acre.

It is evident that the number of acres required to feed one adult varies

TABLE 57

ACRES OF CROP LAND REQUIRED TO PRODUCE DIFFERENT-COST ADEQUATE DIETS,  
THE NUMBER OF PEOPLE THAT CAN BE SUPPORTED WITH EACH DIET, AND THE  
PERCENTAGE OF FOOD ENERGY FROM LIVESTOCK PRODUCTS IN EACH DIET,  
UNITED STATES

| Diet Plan             | Crop Land Required per Person, acres | Number of People that Can Be Supported, millions | Food Energy in Diet from Livestock, per cent |
|-----------------------|--------------------------------------|--|--|
| Low-cost              | 2.12                                 | 203  | 30   |
| Moderate-cost         | 2.57                                 | 167  | 36   |
| Liberal-cost          | 3.15                                 | 137  | 44   |
| 1943-45 civilian diet | 2.67                                 | 161  | 38   |

considerably with the proportion of the diet supplied from livestock products. This is understandable when it is remembered that a given quantity of wheat supplies from 5 to 10 times as many calories when eaten directly as when it is fed to livestock and the meat, lard, butter, or milk is consumed.

The acreage needed to feed one person also varies with the efficiency of production. The total acreage of crop land has not changed much since 1920, although the acreage of crops harvested has increased approximately 3 per cent. In addition, some 55,000,000 acres of land formerly used to grow feed for horses and mules is now being used to produce other crops. On the other hand, food production was about one-third higher in the 1942-45 period than between 1935 and 1939. With advances in science, production can go still higher if there is sufficient incentive to induce the farmer to increase production.

Reference to the number of acres of available land per person given on page 463 shows that we do not have enough good land which may be farmed, without employing soil-conserving methods, to supply even an adequate low-cost diet to our people. But, by employing good management practices on considerable land now being cropped and by bringing into production an additional acreage which must be farmed by soil-conserving methods, there would be some 3.32 acres per person. The data in Table 55 show that few of the countries listed have sufficient arable land to provide even a low-cost diet for their people if a production per acre similar to that in the United States is assumed. In some countries the yield of crops per acre is much higher than in the United States, and in some countries it is very much lower.

**Need for Increased Food Production.** People all over the world are asking for more food. It is doubtful if all the people in any country are adequately fed. In some countries this is not due to a lack of food supplies but to insufficient means to buy enough food, and sometimes to ignorance regarding an adequate diet. On the basis of calories, if it is assumed that about 2,640 calories per person per day are required, the food supplies at the retail level in some 41 countries out of 70 were found inadequate in prewar days. Results from such surveys cannot present an entirely true picture of the food situation because in some instances considerable food is derived from nuts, roots, wild berries, game animals, and similar sources which would not be covered in the survey. On the other hand, deficiencies in minerals, fats, and vitamins exist in diets that furnish a sufficient number of calories.

In considering the food needs in 70 countries for 1960, the food and agriculture organizations of the United Nations have arrived at the data presented in Table 58. According to these estimates a large increase

TABLE 58

ESTIMATED PREWAR PRODUCTION OF BROAD CLASSES OF FOOD AND THE INCREASES AND TARGETS OF THE WORLD FOOD SURVEY \* †

|  | Cereals | Roots ‡ and<br>Tubers | Sugar | Fats<br>and<br>Oils | Pulses<br>and<br>Nuts | Fruits<br>and<br>Vege-<br>tables | Meat § | Milk  |
|--|---------|-----------------------|-------|---------------------|-----------------------|----------------------------------|--------|-------|
| Estimated prewar production, millions of metric tons             | 300.4   | 153.2                 | 30.0  | 15.2                | 36.2                  | 156.3                            | 65.6   | 150.2 |
| Recommended increase to reach 1960 targets, per cent             | 21      | 27                    | 12    | 34                  | 80                    | 163                              | 46     | 100   |
| Targets for world population in 1960,    millions of metric tons | 363.5   | 194.6                 | 33.6  | 20.4                | 65.2                  | 411.0                            | 95.8   | 300.4 |

\* From *Food: Problems and Prospects*, by C. E. Kellogg.

† For 70 countries, including about 90 per cent of the world's population.

‡ Includes bananas.

§ Includes eggs and fish.

|| Assuming a 25 per cent increase in population over 1946.

in all kinds of foods will be needed by 1960. The increases vary from 12 per cent for sugar to 163 per cent for vegetables and fruits. This is a "large order." There are grounds for believing that these targets can be obtained, but it is doubtful that they will be.

**LAND POLICIES OF THE UNITED STATES**

In the early history of our country the people lived to a large extent directly on the products of the soil they tilled. In those days the importance of the farm in the production or creation of wealth, both for the individual and for the political units through taxation, was appreciated to a much greater extent than at present, because industry now provides employment for so many people and has become a fertile source of income to tax-levying agencies. Problems having to do with land accordingly received considerable attention when our country was younger.

In the United States some 30,000,000 people were living on farms in 1930, and another 23,000,000 were classed as rural but non-farm residents. At present not more than 30,000,000 persons are actually on farms, and 34,500,000 are classed as rural but non-farm residents, that is, people living in the country and in towns with less than 2,500 inhabitants but not on farms. This leaves 85,500,000 people classed as urban. Likewise, in many other countries an appreciable proportion of the population lives on the land. In fact, about 70 per cent of the world's population lives on the land. In addition, many persons are engaged in businesses which involve the processing and marketing of farm products and the manufacture and sale of materials used exclusively in connection with farming. As land is so closely connected with the lives and incomes of so many people, it would seem that there should be some definite national policy concerning the ownership, management, and use of land. The question of land policies may be considered under the following headings.

**Questions**

1. Why was the government so anxious to have land pass into private ownership?
2. By what means was ownership of land transferred to individuals and organizations?
3. What attention was paid to quality of land in passing it from public to private ownership?
4. What benefits have resulted from getting land into private ownership?
5. How has misfortune come to people through improper land use?
6. What problems has the land situation brought to public agencies?
7. Are efforts being made to rectify past mistakes?

**Government-Owned Land.** When our government was organized, all land not in private ownership became federal property. The states

and territories held title to no land when they were organized. State ownership of land was a later development by government grant, through purchase, and as a result of tax delinquency. The government was in need of income, and federally owned property brought in little if any returns, whereas privately owned land could be taxed. Farm products were also needed for export so that manufactured goods might be purchased and trade developed with European countries. It was desired also to develop the resources of the country so that the nation might become stronger economically and greater in population. As a result, a great effort was made to get land into private ownership. As the states developed and came into possession of land, they followed the same policy in order to get the land on the tax roll.

**Government Sales and Grants of Land.** Large tracts of land were opened to homesteaders after the Homestead Act in 1862, the areas made available by this method being pushed farther and farther westward as the country became settled. In fact, it was not until the passage of the Taylor Act in 1934 (amended in 1936) that public land ceased to be available to homesteaders. In order to induce railroads to extend their lines westward and to help them finance the construction, the government made railroad companies large grants of land. These lands were then sold to pioneer farmers. A real estate department to dispose of these lands was an important unit in most railroad organizations.

Grants were made to help finance and encourage the development of schools, and thus they hastened the movement of property into private hands. Large acreages in timbered areas were sold to lumber companies, and, where mineral resources were thought to be present, mining companies purchased extensive tracts.

**"Land Was Land."** In the early days there was so much land that little thought was given to the question of soil fertility except by the owners of the older farms near the eastern seaboard, such as Washington and Jefferson. Most land was fairly productive because of its virgin fertility. Anyway, if soil gave out, there was an unlimited supply and so there was no need for concern. Land was opened for homesteading or was sold at extremely low prices without thought of its suitability for farming. The consuming idea was to get people on the land so that it could be placed on the tax roll.

**Advantages of Private Ownership of Land.** The past policy of promoting private ownership of land has resulted in individual benefits to many people and in the rapid production of wealth. The opportunity to obtain land by homesteading or by purchase at low cost has greatly hastened the development of our country. Furthermore, the

possibility of obtaining ownership of land and of passing on that ownership to their children has encouraged industry and initiative and fostered self-reliance, independence, and pride of citizenship in our people. The right to own property is no small part of our democratic liberties.

**Settlers on Poor Land.** The policy of getting land into private ownership with little regard to the quality of the land or its fitness for the proposed use could be expected to result in much misuse of land and consequent human grief and suffering. Thousands of families seeking farm homes and an opportunity to make a living from the soil have found themselves located where successful farming was impossible under conditions which have prevailed. Some of the circumstances which have contributed materially to the unsatisfactory farming conditions are (1) soils of low natural fertility because of limited plant-food content or low moisture-holding capacity or both; (2) a geographical location such that marketing of farm products either was too expensive or required so much time that the products deteriorated in quality; (3) climatic conditions such as short growing season or low temperature during the growing period of crops which limited yields or selection of crops to too great an extent; and (4) rainfall that was too limited or not properly distributed during the growing period for profitable crop production.

In far too many instances large tracts of unproductive land have fallen into the hands of unprincipled operators who have sold farm-sized units to settlers who were not capable of selecting land suitable for farming. The family savings were used as a part payment for the land, with a mortgage on the entire acreage given as security for payment of the balance. When the mortgage was foreclosed, the family was left penniless and the land was resold to another unfortunate settler.

An equally discouraging fate has been experienced by settlers who took up homesteads in areas of low rainfall. A succession of drought years, sometimes accompanied by soil blowing, has necessitated abandonment of many farms. At times the acreage allotment was too small for extensive grazing for which the land was fitted, and the settler was forced to engage in a type of farming unsuitable to his location. The result has been disastrous to both the farmer and the soil.

**Problems Resulting from Poor Land Use.** The inability of farmers, heedlessly located on inferior soils, to make a living has resulted in a great welfare load for local and state agencies dealing with these problems. Schools had to be provided for the children of these families and roads maintained for their convenience. On the other hand, little or no taxes could be collected from them. In some states large acreages

of such lands have reverted to state ownership because of tax delinquency, and the prospects are for a further increase in the amount of state-owned land through this channel. When the land reverts to public ownership, the state is confronted with the problem of the best use for it. This problem is causing much concern, and the question of the adjustment to be made to put these families on a self-supporting basis is yet more urgent. The early policy of getting land into private ownership regardless of the quality of the soil hastened the economic development of the public domain but has ultimately resulted in the return of very large acreages to public ownership, often entailing much human grief, and has left some very serious social and economic problems to be solved.

**Plans for Better Land Use.** Past land policies have resulted in individual benefits and rapid production of wealth, but they have resulted also in waste of soil resources and in a great deal of human distress. As a result of the various and more recent conservation movements, coupled with the economic depression, governmental administrators have become aware of the necessity of correcting the mistakes of the past and of preventing the further serious misuse of land. Various measures for better land use and soil conservation have been devised and are being put into effect.

In order to achieve better land use, basic information about the present extent and distribution of our land resources must be obtained. Land classification and land inventory work are being carried on by such agencies as the National Resources Committee, the U. S. Bureau of Agricultural Economics, the Division of Soil Survey of the U. S. Bureau of Plant Industry, Soils and Agricultural Engineering, the U. S. Soil Conservation Service, and several state experiment stations connected with the land-grant colleges.

Land-planning movements also have been initiated for the purpose of achieving better land use. The ideal objective of rural land planning is to achieve for separate natural and economic land divisions the uses for which they are best adapted under particular local conditions, and then to adjust local plans to state and national plans. Plans which have the sanction of public opinion may result in better land use without legal compulsion, or they may be preliminary to laws regulating use.

Rural zoning ordinances, analogous to city zoning ordinances, have been adopted in a few states. Two of the ostensible purposes of rural zoning are to prevent further agricultural use of submarginal land and to restrict use to that land which is most suitable, thereby preventing

waste in human effort and at the same time protecting land dedicated to non-agricultural uses.

Soil conservation districts may be established by law in some states. The primary purpose of this measure is to restore lands which have deteriorated because of soil erosion and to maintain soil productivity by good management practices. Districts may be any size, and by agreement of landowners or farm operators regulations for land use may be imposed by law. Acquisition, by purchase by national and state agencies, of privately owned poor land and of exploited and so-called submarginal land is another measure proposed for better land use. Lands so acquired may be used for forests, parks, game refuges, public hunting grounds, and other non-agricultural purposes.

Educational programs are also regarded as an essential part of movements for better land use. Extension divisions of land-grant colleges and the U. S. Department of Agriculture establish demonstration projects and arrange meetings and conferences to promote better use of land.

## Glossary

**ACID SOIL.** A soil giving an acid reaction (below *pH* 7.0). A soil having a preponderance of H ions over OH ions in the soil solution.

**ADHESION.** The attraction or union of unlike materials. The particles or substances are attached more or less firmly together by their adjacent surfaces.

**ADSORPTION.** The concentration of material at the contact zone (interface) of two substances. Usually applies to the molecular or ionic state of division; for example, NH<sub>3</sub> gas concentrated on the surface of charcoal, potassium ions concentrated on the surface of soil colloids.

**AGGREGATE (*soil*).** A single mass or cluster of soil particles, such as a clod, crumb, or granule.

**ALKALI SOIL.** A soil containing alkali salts, usually sodium carbonate, or in which the colloidal complex contains a large amount of sodium, with a *pH* value of 8.5 and higher.

**ALKALINE SOIL.** Any soil that is alkaline in reaction, that is, above *pH* 7.0.

**ALLUVIAL SOILS.** Azonal group of soils developed from transported and relatively recently deposited material (by streams) characterized by a weak modification (or none) of the original material by soil-forming processes.

**ALLUVIUM.** Fine material, such as sand, silt, clay, or other sediments deposited on land by streams.

**AMMONIFICATION.** Formation of ammonium compounds or ammonia, as in soils, by soil organisms.

**ANION.** An ion carrying a negative charge of electricity.

**ANTIBIOTICS.** Active chemical substances produced by organisms which are antagonistic to other organisms. Sometimes used to refer specifically to those agents produced by microorganisms that have the capacity of inhibiting the growth of or destroying other microorganisms.

**AZONAL SOILS.** Any group of soils which, owing to their youth or conditions of parent material or relief, are lacking well-developed profile characteristics, and which prevent the development of normal soil-profile characteristics.

**BOG SOILS.** An intrazonal group of soils with a muck or peaty surface soil underlain by peat, developed under swamp or marsh types of vegetation; mostly in a humid or subhumid climate.

**BROWN SOILS.** A zonal group of soils having a brown surface horizon which grades into lighter-colored soil and finally into a layer of carbonate accumulation; developed under short grasses, bunch grasses, and shrubs in a temperate to cool semi-arid climate.

**BUFFERING.** The resistance of a substance to an abrupt change in acidity or alkalinity.

**CALCAREOUS SOIL.** Soil containing sufficient calcium carbonate (often with magnesium carbonate) to effervesce visibly to the naked eye when treated with hydrochloric acid. Soil alkaline in reaction owing to the presence of free calcium carbonate.

**CALICHE.** A more or less cemented deposit of calcium carbonate or of mixed calcium and magnesium carbonates, characteristic of soils of warm or hot desert and semi-arid regions.

**CAPILLARY CAPACITY (*field capacity*).** The amount of water held in the soil after the excess or gravitational water has drained away.

**CAPILLARY WATER.** Water held by adhesion and surface-tension forces as a film around particles and in the capillary spaces. Moves in any direction in which capillary tension is greatest.

**CARBON-NITROGEN RATIO.** The relative proportion, by weight, of organic carbon to nitrogen in a soil. The number obtained by dividing the percentage of organic carbon in a soil by the percentage of nitrogen.

**CARBONATE ACCUMULATION, SOIL HORIZON OF.** A developed soil horizon, beneath the surface, containing more calcium (or magnesium and calcium) carbonate than the soil above it or the soil material below it.

**CATCH CROP.** A crop seeded with one of the regular crops in a rotation or between the growing periods of two regular crops, for the purpose of adding organic matter and, in some cases, nitrogen also to the soil.

**CATEGORY (*soil classification*).** Any one of the subdivisions of the system of classification in which soils are arranged on the basis of their characteristics. Beginning with the lowest category, soils are classified on the basis of progressively fewer characteristics into groups of progressively higher or more inclusive categories.

**CATION.** An ion carrying a positive charge of electricity.

**CATION EXCHANGE.** See **ION-EXCHANGE**.

**CATION-EXCHANGE CAPACITY.** The maximum quantity of cation that the soil is capable of adsorbing from a neutral solution of a monovalent cation. Expressed in terms of milliequivalents per 100 grams of soil.

**CHERNOZEM SOILS.** A zonal group of soils having a deep, dark-colored to nearly black surface horizon, rich in organic matter, which grades below into lighter-colored soil and finally into a layer of lime accumulation; developed under tall and mixed grasses in a temperate to cool subhumid climate. From the Russian for black earth. Sometimes spelled **TSCHERNOSEM**.

**CHESTNUT SOILS.** A zonal group of soils having a dark brown surface horizon which grades below into lighter-colored soil and finally into a horizon of lime accumulation; developed under mixed tall and short grasses in a temperate to cool and subhumid to semi-arid climate. They occur on the arid side of Chernozem soils, into which they grade.

**CLASS, SOIL.** Classes of soil based on the relative proportion of soil separates.  
**CLAY.** The small mineral soil grains or particles, less than 0.002 mm. in diameter.

**CLAYPAN.** A dense and heavy soil horizon underlying the upper part of the soil; hard when dry and plastic or stiff when wet; presumably formed in part by the accumulation of clay brought in from the horizons above by percolating water.

**COHESION.** The union or attraction of substances of like characteristics, as the attraction of one water molecule for another.

**COLLOID, SOIL.** The term colloid is used in reference to matter, both inorganic and organic, having very small particle size and a correspondingly high surface area per unit of mass. Many mineral colloids exhibit crystalline structure. Colloid comes from the Greek words for glue and like.

**COLLUVIA.** Heterogeneous deposits of rock fragments and soil material accumulated at the base of comparatively steep slopes through the influence of gravity.

**COVER CROP.** A crop used to cover the soil surface; to decrease erosion and leaching, shade the ground, and offer protection to the ground from excessive freezing and heaving.

**DEFLOCULATE.** To separate or break down soil aggregates into their component particles. Usually refers to particles of colloidal dimensions.

**DENITRIFICATION.** The reduction of nitrates to nitrites, ammonia, and free nitrogen, as in soil by soil organisms, particularly certain anaerobic organisms (those living or active in the absence of air or free oxygen).

**DISPERSION.** The destroying of soil structure (breaking up the granules) so that each individual soil particle behaves as a unit.

**DRIFT.** Material of any sort deposited in one place after having been moved from another. For example, glacial drift includes glacial deposits, unstratified (till) and stratified glacial outwash materials.

**DRUMLIN.** An oval hill of glacial drift, normally compact and unstratified, usually with its longer axis parallel to the movement of the ice responsible for its deposition.

**ELUVIATION.** The movement of soil material from one place to another within the soil, in solution or in suspension, where there is an excess of rainfall over evaporation. Horizons that have lost material through eluviation are referred to as eluvial and those that have received material as illuvial.

**EROSION (*geological or normal*).** Erosion suffered by land in its natural state undisturbed by human activity. Erosion taking place on land used for crop production or on land whose natural condition has otherwise been modified through man's activities is sometimes referred to as "soil erosion."

**EROSION (*land*).** The wearing away of the land surface by falling or running water, by wind, or by other natural agents such as gravity.

**FAMILY, SOIL.** A category in soil classification between series and great soil group; a taxonomic group of soils having similar profiles, composed of one or more distinct soil series.

**FERTILITY (*of soil*).** The quality that enables a soil to provide the proper compounds, in the proper amounts and in the proper balance, for the growth of specified plants when other factors, such as light, temperature, and the physical condition of the soil, are favorable.

**FERTILIZER.** A material supplying one or more of the plant nutrients in a condition suitable for application to the soil.

**FLOCCULATE.** To aggregate individual particles into small groups or clusters. Usually refers to particles of colloidal dimensions.

**FLOOD PLAIN.** The nearly flat surface subject to overflow along streams.

**FRIABLE.** Easily crumbled in the fingers; non-plastic.

**GENESIS, SOIL.** Mode of origin of the soil; refers particularly to the processes responsible for the development of the solum from the unconsolidated parent material.

**GEY.** A soil horizon in which the material is usually bluish gray or olive gray, more or less sticky, compact, frequently without definite structure; developed under the influence of excessive moisture.

**GRANULATION.** The cementation of particles into masses as grains, aggregates, or clumps, essentially a result of flocculation and aggregation.

**GRAVITATIONAL WATER.** The water that moves under the force of gravity; it is not retained by the soil.

**GRAY-BROWN PODZOLIC SOILS.** A zonal group of soils having a comparatively thin organic covering and organic-mineral layers over a grayish brown leached layer which rests upon an illuvial brown horizon; developed under deciduous forest in a temperate moist climate.

**GREAT SOIL GROUPS (*soil classification*).** A group of soils having common internal soil characteristics; includes one or more families of soils.

**GREEN-MANURE CROP.** Any crop grown and worked into the soil for the purpose of improving the soil, especially by the addition of organic matter.

**HARDPAN.** A hardened or cemented soil horizon. The soil may have any texture and is compacted or cemented by iron oxide, organic material, silica, calcium carbonate, or other substances.

**HORIZON, SOIL.** A layer of soil approximately parallel to the land surface with more or less well-defined characteristics that have been produced through the operation of soil-building processes. Each layer differs from the one above or below in some characteristic.

**HUMUS.** The well-decomposed, more or less stable part of the organic matter of the soil.

**HYDROLYSIS.** A double decomposition reaction in which water is a reactant.

**HYGROSCOPIC COEFFICIENT.** The maximum amount of water (percentage based on weight of dry soil) adsorbed on the surface of soil particles from an atmosphere slightly below 100 per cent relative humidity.

**HYGROSCOPIC WATER.** Water which is adsorbed from atmospheric water vapor and held on the surface of particles by forces of adhesion.

**IGNEOUS ROCK.** A rock produced through the cooling of melted mineral material.

**ILLUVIAL HORIZON.** A soil horizon that has received material from other portions of the soil profile through the process of eluviation.

**IMMATURE SOIL.** A young or imperfectly developed soil.

**INFILTRATION.** The process by which water enters the soil through the surface. The rate at which water soaks into the soil is called the "infiltration capacity."

**INTRAZONAL SOIL.** Any of the great groups of soils with more or less well-developed soil characteristics that reflect the dominating influence of some local factor of relief, parent material, or age over the normal effect of the climate and vegetation.

**ION.** An electrically charged element or group of elements in an electrolyte. An electrically charged particle.

**ION-EXCHANGE.** The replacement in a colloidal system of an ion by another with a charge of the same sign. Cation exchange refers to the interchange of positively charged ions (cations), and anion exchange refers to the replacement of negatively charged ions (anions) by other anions.

**LACUSTRINE DEPOSITS.** Materials deposited by lake waters.

**LAND, ARABLE.** Land which, in its present condition, is physically capable, without further substantial improvement, of producing crops requiring tillage.

**LAND RECLAMATION.** Making land capable of more intensive use by changing its character or environment or both through operations requiring collective effort. Does not include the clearing of stumps, brush, and stones from land, or simple techniques of erosion control that can be effected by the individual.

**LAND TYPE.** Land uniformly possessed of particular distinguishing characteristics. A natural land type is land having a particular set of defined natural characteristics, principally of soil, climate, relief, stoniness, and native vegetation.

**LAND-USE PLANNING.** A development of plans for the uses of land that will, over a long period, best serve the general welfare, together with the formulation of ways and means of achieving such uses.

**LANDSCAPE** (*as used in soil geography*). The sum total of the characteristics that distinguish a certain area on the earth's surface from other areas. These characteristics are the result not only of natural forces but of human occupancy and use of the land.

**LATERITE SOILS.** It has been recommended that the use of this term be discontinued as defined in "Soils and Men." The term "latosol" has been suggested as a replacement to refer to the more friable and iron-rich soils of the hot, moist, or wet-dry tropics referred to as "red loams" in European, Asiatic, and Australian literature.

**LATERIZATION.** The processes by which latosols are produced; that is, soils having the following characteristics: (a) a low silica-sesquioxide ratio in the clay fraction, (b) low activity of clay with medium to low cation-exchange capacity of the mineral fraction, (c) low content of all but the most resistant primary minerals, (d) low content of soluble material, (e) high degree of aggregate stability, and (f) red color or reddish shades of other colors. Horizons of concentration of iron and aluminum oxides may be present as a result of removal of silica.

**LEACHING.** Removal of materials in solution.

**LIME, AGRICULTURAL.** Any compound of calcium or magnesium or both used to correct the harmful effects of acid soils on plant growth.

**LIMESTONE.** A general name for rocks composed essentially of calcium carbonate. There are several kinds of limestone, and they vary in physical and chemical composition.

**LITHOSOLS.** An azonal group of soils having no clearly expressed soil morphology and consisting of a mass of rock fragments from consolidated rocks which are imperfectly weathered. They are found primarily on steeply sloping land. They have only recently been differentiated from regosols.

**LOESS.** Wind-transported material of fine texture.

**MARINE MATERIAL.** Material deposited in the waters of oceans and seas and exposed by elevation of the land or the lowering of the water level.

**MARL.** A soft, earthy deposit consisting chiefly of calcium carbonate mixed with sand, clay, organic matter, and other impurities in varying proportions. Frequently used as a liming material.

**MATURE SOIL.** A soil with well-developed characteristics, produced by the natural processes of soil formation, and in equilibrium with its environment.

**MECHANICAL ANALYSIS.** The separation by mechanical means of the different size groups (separates) and the determination of the percentage of each group in a given soil sample.

**METAMORPHIC ROCK.** A rock the constitution of which has undergone pronounced alteration. Such changes are generally effected by the

combined action of pressure, heat, and water, frequently resulting in a more compact and more highly crystalline condition of the rock. Gneiss, schist, and marble are common examples.

**MINERAL.** A naturally occurring combination of inorganic elements and ions in the form of salts or other compounds either in crystalline or amorphous condition.

**MINERAL SOIL.** A general term used in reference to any soil composed chiefly of mineral matter. The mineral material is dominant over the organic matter in determining the characteristics of the soil.

**MOISTURE EQUIVALENT.** The amount of water retained (percentage) by a soil against a force of 1,000 times that of gravity.

**MORPHOLOGY, SOIL.** The physical constitution of the soil, including the texture, structure, porosity, consistence, and color of the various soil horizons, their thickness, and their arrangement in the soil profile.

**MOTTLED (*mottling*).** Irregularly marked with spots of different colors.

**MUCK SOIL.** An organic soil, composed of highly decomposed organic material, with a considerable amount of mineral soil material, finely divided and with few fibrous remains of the original plants.

**NEUTRAL SOIL.** A soil that is not significantly acid or alkaline; strictly, one having a pH of 7.0.

**NITRIFICATION.** Formation of nitrates from ammonia as in soils by soil organisms.

**NITROGEN FIXATION.** In soils, the assimilation of free nitrogen from the air by soil organisms, making the nitrogen eventually available to plants.

**NORMAL SOIL.** A soil having a profile in equilibrium with the two principal forces of the environment—native vegetation and climate—usually developed on the gently undulating upland; with good drainage; from any parent material; not of extreme texture or chemical composition; that has been in place long enough for biological forces to exert their full effect.

**NUTRIENTS, PLANT.** The elements or groups of elements taken in by the plant, essential to its growth, and used by it in the elaboration of its food and tissue.

**ORGANIC SOILS.** Soils containing organic matter in sufficient quantities to dominate the soil characteristics. Frequently all soils containing 20 per cent or more organic matter by weight are arbitrarily designated as organic soils.

**OXIDATION.** The process that takes place when an element burns in oxygen; the product is a compound called an oxide and the process is called oxidation. In a broader sense, when the positive valence of an element is increased, that element is oxidized.

**PARENT MATERIAL.** The slightly altered or unweathered material beneath the solum; similar to that from which the soil was formed.

**PARENT ROCK.** The rock from which parent materials of soil are formed.

**PEAT.** Unconsolidated soil material consisting primarily of undecomposed or slightly decomposed organic matter accumulated under conditions of excessive moisture.

**PEDALFER.** A soil in which there is a zone of alumina and iron oxide accumulation in the profile but with no horizon of carbonate accumulation.

**PEDOCAL.** A soil with a horizon of accumulated carbonates in the soil profile.

**PEDOLOGY.** The science of the soil in which different soils are considered natural units and attention is given to development, physical, chemical, and biological relationships, and to their dynamic nature.

**PERCOLATION.** The process whereby water moves through the soil in response to the force of gravity.

**pF VALUE.** The logarithm of the height, in centimeters, of a water column necessary to produce a force equal to the energy with which moisture is held by a soil. The *p* indicates a logarithm, and the F is used to suggest energy.

**pH.** A notation to designate or indicate the degree of acidity or alkalinity of systems. Technically, the common logarithm of the reciprocal of the hydrogen-ion concentration (grams per liter) of a system.

**PHASE, SOIL.** That part of a soil type having minor variations in characteristics used in soil classification from the characteristics normal for the type, although they may be of great practical importance. The variations are chiefly in such external characteristics as relief, stoniness, or erosion.

**PLANOSOL.** An intrazonal group of soils with leached surface horizons underlain by strongly contrasting *B* horizons which are more strongly illuviated, cemented, or compacted than those in normal soils. They are developed in humid or subhumid climates and under grass or forest cover. It has been suggested that planosols be elevated to the rank of a suborder of intrazonal soils and the concept of them be broadened to include Ground Water Podzols, Ground Water Laterites, and poorly drained soils having "silt pans," "hardpans," or "claypans."

**PLASTIC.** A soil which can be readily molded or deformed without rupture; pliable, putty-like.

**PODZOL SOILS.** A zonal group of soils having an organic mat and a very thin organic-mineral layer above a gray leached layer which rests upon an illuvial dark brown horizon; developed under the coniferous or mixed forest or under heath vegetation in a temperate to cold moist climate. Iron oxide and alumina, and sometimes organic matter, have been removed from the *A* and deposited in the *B* horizon.

**PODZOLIC SOILS.** Soils that have been formed wholly or partly under the influence of the podzolization process.

**PODZOLIZATION.** A general term referring to that process (or those processes) by which soils are depleted of bases, become acid, and have developed eluvial *A* horizons (surface layers of removal) and illuvial *B* horizons (lower horizons of accumulation). Specifically the term refers to the process by which a podzol is developed, including the more rapid removal of iron and alumina than of silica, from the surface horizons, but it is also used to include similar processes operative in the formation of certain other soils of humid regions.

**POROSITY, SOIL.** The degree to which the soil mass is permeated with pores or cavities. It is expressed as the percentage of the whole volume of the soil which is unoccupied by solid particles.

**PRAIRIE "BRUNIGRA" SOILS.** The zonal group of soils having a very dark brown or grayish brown surface horizon, grading through brown soil to the lighter-colored parent material at 2 to 5 feet, developed under tall grasses, in a temperate, relatively humid climate. This term is not applied to all dark-colored soils of the treeless plains but only to those in which carbonates have not been concentrated in any part of the profile by the soil-forming processes.

**PRODUCTIVITY (*of soil*).** The capability of a soil for producing a specified plant or sequence of plants under a specified system of management.

**PROFILE, SOIL.** A vertical cross section of the soil from the surface into the underlying unweathered material.

**PUDDLE.** To deflocculate or to destroy the granular structure of the soil.

**REACTION, SOIL.** The degree of acidity or alkalinity of the soil mass expressed in pH values or in words.

**REDUCTION.** The process by which an element loses valence, that is, a decrease in positive valence or an increase in negative valence. Often used to designate the loss of oxygen from a compound.

**REGOSOLS.** An azonal group of soils consisting of deep soft mineral deposits (unconsolidated rock) in which few if any clearly expressed soil characteristics have developed; largely confined to recent sand dunes, loess, and glacial drift of steeply sloping lands.

**RELIEF.** The elevations or inequalities of a land surface considered collectively.

**RESIDUAL SOIL.** Soils formed by the weathering of rock and minerals in place; sedentary materials.

**ROCK.** Any relatively homogeneous mass of mineral material that forms a portion of the earth's crust, including loose, incoherent masses as well as solid masses. The common concept that material must be hard, as granite or limestone, to be classed as rock is not correct from a geological point of view.

**SALINE SOIL.** A soil containing excessive amounts of neutral or non-alkaline salts, usually chlorides and sulfates.

**SALT.** The product, other than water, of the reaction of a base with an acid.

**SAND.** Small rock or mineral fragments having diameters ranging from 1 to 0.05 mm.

**SEDENTARY SOILS.** Soils formed in place without the addition of transported material.

**SEDIMENTARY ROCK.** A rock composed of particles deposited from suspension in water.

**SEPARATE, SOIL.** One of several groups of soil particles having definite size limits.

**SERIES, SOIL.** A group of soils having genetic horizons similar in differentiating characteristics and arrangement in the soil profile, except for the texture of the surface soil, and developed from a particular type of parent material. A series may include two or more soil types differing from one another in the texture of the surface soils.

**SILT.** Small mineral soil grains the particles of which range in diameter from 0.05 to 0.002 mm. (or 0.02 to 0.002 mm. in the international system).

**SOIL.** The natural medium for the growth of land plants on the surface of the earth. A natural body on the surface of the earth in which plants grow; composed of organic and mineral materials.

**SOLONCHAK.** An intrazonal group of soils having a high concentration of soluble salts; usually light colored; without characteristic structural form; developed under salt-loving grass or shrub vegetation mostly in an arid, semi-arid, or subhumid climate.

**SOLONETZ SOILS.** An intrazonal group of soils having a variable surface horizon of friable soil underlain by dark hard soil, ordinarily with columnar structure; usually highly alkaline; developed under grass or shrub vegetation, mostly in a subhumid or semi-arid climate.

**SOLOTH SOILS.** An intrazonal group of soils having a thin surface layer of brown, friable soil over a gray leached horizon resting on a brown or dark brown horizon. They develop in a semi-arid or subhumid climate under shrubs, grasses, or a mixture of grass and trees.

**SOLUM.** The upper part of the soil profile, above the parent material, in which the processes of soil formation are taking place. In mature soils this includes the *A* and *B* horizons, and the character of the material may be and usually is greatly unlike that of the parent material beneath. Living roots and life processes are largely confined to the solum.

**SPECIFIC GRAVITY.** The ratio of the weight of dry soil to that of water which will occupy only the volume of the soil particles alone (pore space excluded). It is the ratio between the weight of an object and the weight of water it will displace.

**STRATIFIED.** Composed of, or arranged in, strata or layers, as stratified alluvium. The term is applied to geological materials. Those layers in soils that are produced by the processes of soil formation are called horizons; those inherited from the parent material are called strata.

**STRIP CROPPING.** Strip cropping is a practice of growing ordinary farm crops in long strips of variable widths, across the line of slope, approximately on the contour, in which dense-growing crops are seeded in alternate strips with clean-tilled crops.

**STRUCTURE, SOIL.** The morphological aggregates in which the individual soil particles are arranged.

**SUBSOIL.** Roughly, that part of the solum below plow depth.

**SULFOFICATION.** The oxidation of sulfur, either free or from organic or inorganic compounds, into oxides of sulfur largely through the action of microorganisms. In the soil these oxides are quickly changed into sulfites and sulfates.

**SURFACE SOIL.** That part of the upper soil of arable soils commonly stirred by tillage implements or an equivalent (5 to 8 inches) in non-arable soils.

**TERRACE** (*for control of runoff, soil erosion, or both*). A broad surface channel or embankment constructed across the sloping lands, on or approximately on contour lines, at specific intervals. The terrace intercepts surplus runoff to retard it for infiltration or to direct the flow to an outlet at non-erosive velocity.

**TEXTURE, SOIL.** The relative proportion of the various size groups of individual soil grains. The coarseness or fineness of the soil.

**TILL, GLACIAL.** A deposit of earth, sand, gravel, and boulders transported by glaciers. Till is unstratified.

**TILL PLAIN.** A level or undulating land surface covered by glacial till.

**TILTH, SOIL.** The physical condition of the soil in relation to plant growth. A term indicating the conditions of soil structure produced by tillage or cultivation.

**TOP SOIL.** A general term applied to the surface portion of the soil, including the average plow depth (surface soil) or the *A* horizon, where this is deeper than plow depth.

**TUNDRA SOILS.** A zonal group of soils having dark brown highly organic layers over grayish horizons which rest on an ever-frozen substratum; developed under shrubs and mosses in cold, semi-arid to humid climates, that is, in Arctic regions.

**TYPE, SOIL.** A group of soils having genetic horizons similar in differentiating characteristics, including texture and arrangement in the soil profile, and developed from a particular type of parent material.

**VARNISH, DESERT.** A glossy coating of dark-colored compounds, probably composed mainly of iron oxides, covering pebbles, stones, and large rock surfaces exposed in hot deserts.

**VESICULAR STRUCTURE.** Soil structure characterized by round or egg-shaped cavities or vesicles.

**VOLUME WEIGHT.** The weight of a given volume of dry soil, in its natural structural conditions, in comparison to the weight of an equal volume of water. It is sometimes referred to as the *apparent specific gravity*.

**WATER REQUIREMENT OF PLANTS** (*transpiration ratio*). The pounds of water transpired by a plant per pound of dry matter produced above ground.

**WATER TABLE.** The upper limit of the part of the soil or underlying material wholly saturated with water.

**WEATHERING.** The physical and chemical disintegration or decomposition of rocks and minerals under natural conditions.

**WILTING COEFFICIENT** (*wilting point*). The percentage of water in the soil (based on dry weight of the soil) when permanent wilting of plants occurs. It refers to that moisture content at which soil cannot supply water at a rate sufficient to maintain the turgor of a plant and it permanently wilts.

**ZONAL SOIL.** Any one of the great groups of soils having well-developed soil characteristics that reflect the influence of the active factors of soil genesis—climate and living organisms, chiefly vegetation.

*In the compilation of the Glossary extensive use was  
made of the glossary in "Soils and Men,"  
U.S.D.A. Yearbook, 1938.*

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